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(54) Title: FUEL CELL SYSTEM FOR LOW PRESSURE OPERATION

(57) Abstract

A fuel cell design for use at low pressure. The invention has a reduced number of component parts to reduce fabrication costs, as well as a simpler design that permits the size of the system to be reduced at the same time as performence is being improved. In the present design, an adjacent anode and cathode pair are fabricated using a common conductive element, with that conductive element excited to conduct the current from one cell to the adjacent one. This produces a small and simple system suitable for operating with pass serving to conduct the current from one cell to the adjacent one. This produces a small and simple system suitable for operating with pass are related to the state of the size of the

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FUEL CELL SYSTEM FOR LOW PRESSURE OPERATION

This application is a continuation-in-part application of co-pending U.S. patent application serial no. 08/926,547 filed September 10, 1997.

Field of the Invention

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This invention relates generally to the field of fuel cells. More particularly, the invention relates to low pressure fuel cells and their components.

Background of the Invention

Generally, a fuel cell is a device which converts the energy of a chemical reaction into electricity. Fuel cells differ from batteries in that fuel and oxidant are stored external to the cell, which can generate power as long as the fuel and oxidant are supplied. A fuel cell produces an electromotive force by bringing the fuel and oxidant in contact with two suitable electrodes separated by an electrolyte. A fuel, such as hydrogen gas, is introduced at one electrode where it dissociates on the electrocatalytic surface of the positive electrode (anode) to form protons and electrons, as elucidated in equation 1. The electrons pass into the conductive structure of the electrode, and there from to the external electrical circuit energized by said fuel cell. The protons formed by dissociation of the hydrogen at the first electrode and pass through the electrolyte to the second electrode. Simultaneously, an oxidant, such as oxygen gas or air, is introduced to the second electrode where it is adsorbed on the electrocatalytic surface of the negative electrode (cathode) and is electrochemically reduced to form a surface oxide species by electrons having transversed the external electrical circuit energized by the fuel cell. This surface oxide reacts with protons from the electrolyte to form water, the product of the net reaction. The water desorbs from the electrode and leaves the cell in the cathode gas stream. The half cell reactions for a hydrogen consuming fuel cell at the two electrodes are, respectively, as follows

$$H_2 \rightarrow 2H^+ + 2e^-$$
 (1)

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$$\frac{1}{2}O_{2} + 2H^{+} + 2e^{-} \rightarrow H_{2}O$$
 (2)

Connecting the two electrodes through an extendal circuit causes an electrical current to flow in the circuit and withdraws electrical power from the cell. The overall fuel cell reaction, which is the sum of the separate half cell reactions written above, produces electrical energy and heat.

In practice, fuel cells are not operated as single units, but are connected in a series to additively combine the individual cell potentials and achieve a greater, and more useful, potential. The cells in a given series can be connected directly, with opposing faces of a single component in contact with the anode of one cell and the cathode of an adjacent cell, or through an external electrical linkage. A series of fuel cells, referred to as a fuel cell stack, are normally equipped with a manifold system for the distribution of two gases. The fuel and oxidant are directed with manifolds to the correct electrodes, and cooling is provided either by the reactants or by a cooling medium. Also within the stack are current collectors, cell-to-cell seals, and other components. The stack and associated hardware make up the fuel cell module.

In fuel cells where a solid polymer electrolyte or proton exchange membrane ("PEM") is used, the membrane acts as the electrolyte as well as a barrier for preventing the mixing of the reactant gases. A PEM fuel cell is described in greater detail in Dhar, U.S. Patent No. 5,242,764, which is incorporated herein by reference. Much research and development has been devoted to improving the power-to-weight ratio for proton exchange membrane (PEM) fuel cells. Most of this research has involved increasing the power per unit volume of relatively heavy stacks.

Figure 1 is a drawing illustrating a fuel cell stack based on a conventional bipolar filter press design 10 with graphite structure elements. A full description of filter press type fuel cells may be found in Neidrach, U.S. Patent No. 3,134,697 which is incorporated herein by reference. While improvements in the filter press style fuel cells have provided significant increases in power per unit volume, the overall systems that have evolved are large, heavy, and relatively complex, with compressors to supply air and pumps to provide forced water cooling systems to remove excess heat.

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Figure 2 shows the structure of a standard fuel cell membrane and electrode (M&E) assembly 20 intended for use in the bipolar stack 10 of Figure 1, which has current collection over most of the back of the electrode. The M&E assembly consists of a membrane 22, a catalyst layer 24, a gas diffusion layer 26 and a conductive cloth backing 28. As illustrated, a complete M&E assembly includes similar layers formed on both sides of the membrane.

More recently, efforts have been made to reduce the stack weight by replacing the heavy carbon elements with thinner and lighter, metal elements. However, these units were designed for large scale applications, some on the order of about 30 kW, and, therefore, require the same stack ancillary equipment mentioned above. Furthermore, the ancillary equipment included with the stack in these systems has been designed to operate efficiently at the kilowatt level. Scaled down versions of these systems have been attempted in applications that require much less power, such as within the range between about 50 and about 150 Watts. However, these systems are not well suited for stack outputs in the tens or hundreds of watts, since the rotating components, such as pumps and compressors, do not scale down well. As a result, even small scale systems of this design are too heavy for many small applications, such as for portable applications and personal use.

Therefore, perhaps the most important objective for portable and personal applications is not Watts per unit volume but Watts per unit weight, i.e. W/lb. Efforts to adapt the standard bipolar filter press design to low pressure operation, thereby eliminating much of the ancillary equipment, have met with some limited success, producing stacks with power densities as high as 61 W/lb. While this is a useful power density, these systems require complicated and expensive assembly.

One possible way of improving fuel cell systems for operation at lower pressures is using liquid fuels in lieu of gaseous fuels, such as hydrogen. Methanol (CH₃OH) and other related compounds, such as dimethoxymethane ($C_3H_5O_2$) and trimethoxymethane ($C_4H_{10}O_3$), offer very promising alternatives to gaseous fuels.

The liquid fuels mentioned above share some common advantages compared to hydrogen. First, they are all pourable liquids at ambient pressure and ambient, and near ambient, temperatures. Second, they have a much higher energy density than

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hydrogen. For example, a 1.1 methanol water mixture (each mole of methanol requires a mole of water for electrochemical oxidation, as shown in Equation 3) has as much potential energy as hydrogen stored at a pressure of 16,000 psi.

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$$CH_3OH + H_2O + \frac{3}{2}O_2 \rightarrow CO_2 + 3H_2O$$
 (3)

Although there is a net production of water, e.g., more water is produced at the cathode than consumed at the anode, water must be supplied to the anode because in a fuel cell the oxidation is carried out as a pair of half cell reactions, with Equation 3 representing the net reaction. In a fuel cell water is consumed in the anode reaction (Equation 4) and produced in the cathode reaction (Equation 5).

$$CH_3OH + H_2O \rightarrow CO_2 + 6 H^+ + 6 e^-$$
 (4)

$$6 \text{ H}^+ + 6 \text{ e}^- + \frac{3}{2} \text{ O}_2 \rightarrow 3 \text{ H}_2 \text{O}$$
 (5)

The reactions for dimethoxymethane and trimethoxymethane are similar, with four and five molecules of water needed for each molecule of the oxidized organic compound respectively. Since they are consumed at higher water-to-fuel stoichiometries, the two ethers should be present at lower concentrations in the fuel stream, and consequently have lower permeation rates through PEM membranes and lower vapor pressures over the solution. This is especially true in the case of trimethoxymethane, which has a boiling point of 104 °C, 40 °C higher than methanol, and consequently, a lower vapor pressure than methanol under all conditions. The lower vapor pressure, combined with lower toxicity for these compounds, leads to less vapor toxicity hazards compared to those associated with the use of methanol.

Therefore, there is a need for a lightweight fuel cell system that provides an improved power density (W/lb) and eliminates much of the ancillary equipment. There is also a need for fuel cells that operate on gaseous fuels, such as hydrogen, and fuel cells that operate on liquid fuels, such as dimethoxymethane. It would be desirable if

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the fuel cell operated efficiently in the 50 to 150 Watt range to supply electricity to a variety of common electrical devices. It would also be desirable if the fuel cell had no more than a few moving parts to reduce maintenance and avoid breakdowns. It would be further desirable to have a fuel cell system that was available in modules that could be configured together to meet the power requirements of specific applications.

Summary of the Invention

The present invention provides an apparatus comprising an array of cells. Each cell comprises a membrane, an anode comprising a conductive element, and a cathode comprising a conductive element. The cathode of one cell shares its conductive element with the anode of an adjacent cell; the sharing of conductive elements establishing an electrical connection between the cells in the array. In one embodiment the cells are electrochemical cells suitable for the direct production of electricity through the simultaneous oxidation of a fuel, or reducing agent, at one electrode, the oxide, accompanied by simultaneous reduction of an oxidizer at the other electrode, the cathode. In another embodiment, the cells can be used to produce oxidized and/or reduced products through simultaneous electrochemical oxidation at one electrode and reduction at the opposing electrode of a compound or a mixture of compounds by application of an electrical current. Further, the invention provides for the use of both gas and liquid feeds.

The invention also encompasses a low pressure gas electrode comprising (a) an electrically conducting supporting substrate, wherein said substrate comprises at least two opposite faces; a plurality of openings allowing passage from one face to an opposite face; said openings encompassing between about 10% and about 90 % of the total area of the substrate; and (b) an electrically conductive porous material bound to the substrate comprised of a conductive component and a binding component. In one embodiment, the conductive porous material is a mixture comprising a powdered component and a fibrous component. In yet another embodiment, the powdered and/or fibrous components comprise carbon.

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Brief Description of the Drawings

So that the above recited features and advantages the present invention can be understood in detail, a more particular description of the invention, briefly summarized above, may be had by reference to the embodiments thereof which are illustrated in the appended drawings. It is to be noted, however, that the appended drawings illustrate only typical embodiments of this invention and are therefore not to be considered limiting of its scope, for the invention may admit to other equally effective embodiments.

Figure 1 is a standard filter press type fuel cell stack 10 showing the arrangement of the bipolar cell plates and end plates.

Figure 2 is a cross-sectional view of a standard PEM fuel cell membrane and electrode assembly (M&E) 20.

Figure 3 illustrates the key components for a monopolar cell stack, including a terminal cell frame 36, a flat bipolar cell frame 30 and a curved bipolar cell frame 38.

Figure 4 is a cross-sectional view of a fuel cell stack 50 comprised of monopolar cells based on the use of the bipolar cell frame components of Figure 3.

Figure 5 is a graph showing the effect of both increased fuel pressure and forced convection on the performance of a four cell fuel cell stack.

Figures 6(a-c) are cross-sectional views of three configurations for an M&E produced with a metal conductive element (grid) 82 included as part of the structure.

Figure 7 is a graph showing the improvement in performance produced by attaching a metal grid to the back of the electrode.

Figure 8 is an exemplary monopolar fuel cell stack 95.

Figure 9 is a perspective view of a compact fuel cell 100 having a fan 106 to provide forced convention of air over the cathodes 108.

Figure 10 is an illustration of the components of a monopolar fuel cell constructed without cell frames.

Figure 11 is an expanded metal sheet suitable for use as the conductive element in the electrodes

30 Figure 12(a) is a face on view into the cathodes on one face of a fin.

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Figure 12(b) is a cross-sectional view cut vertically near the center of Figure 12(a).

Figure 12(c) is a view of an enlarged piece cut out of the side of 12(b) indicating the location of the wicking structure.

Figure 13(a) is a drawing showing a wrap-around fuel cell stack supported on its fuel supply, illustrated here as a cylinder with series of square window panes on the surface of the cylinder.

Figure 13(b) is a drawing showing a wrap-around fuel cell stack supported on its fuel supply, illustrated here as a cylinder with a series of bands, with long, narrow individual cells.

Figure 14 is a cross-sectional view of a monopolar fuel cell mounted on the outside of the vessel that supplies its fuel.

Figure 15 is a graph showing polarization curves for M&E's produced using three different conductive elements in the electrodes with the same composition for the gas diffusion layer and the catalyst layer.

Figure 16 is a graph showing polarization curves for M&E's produced using two different membrane materials with the same conductive element (Cu 4/3 in Table II) and the same compositions for both the gas diffusion layer and the catalyst layer.

Figure 17 is an illustration of the arrangement used to focus the press load on one cell of a multi-cell module.

Figure 18 is a cross-sectional view of an electrode fabricated using gas diffusion electrodes with a metal grid as part of the electrode for improved internal conductivity.

Figure 19 is graph showing the polarization curve for a two-cell monopolar module with the cells connected through their common electrode support.

Figure 20 is a graph showing polarization curves for a three-cell monopolar fuel cell module illustrating the effect of fuel humidification on the unit's performance.

Figure 21 is a graph showing the power output of the three-cell monopolar fuel cell module shown operating without humidification in Figure 20.

Figure 22 is a graph showing polarization curves for a three-cell monopolar fuel cell module illustrating the effect of hydrogen pressure on the unit's performance.

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Figure 23 is a graph showing polarization curves for a three-cell monopolar fuel cell module illustrating the effect of air motion on the unit's performance.

Figure 24 is a graph showing the polarization curves for a stack operated with and without air flow from a fan.

Figure 25 is a schematic diagram depicting a polymeric frame.

Figure 26(a) is a cross sectional view of a segment of a fin similar to the fin of Figure 31(b) with an internal heat transfer structure added.

Figure 26(b) is an edge view of the fin of Figure 26(a) showing the internal heat structure.

Figure 27 is a drawing depicting a wavy convoluted aluminum sheet as an example of internal heat transfer structures.

Figure 28 is a cross sectional view of the fin of Figure 26(a) with the back plane of the monopolar cells extended and folded over to form a protective cover.

Figure 29 is a cross sectional view of a fin with protective cooling fins.

Figures 30(a-c) are cross-sectional views of three configurations for an M&E produced with a metal conductive element included as part of the structure.

Figure 31(a) is a face on view into the cathodes on one face of a fin.

Figure 31(b) is a cross-sectional view cut vertically near the center of Figure 21(a).

Figure 31(c) is a cross-sectional view cut horizontally near the center of Figure 31(a).

Detailed Description of the Preferred Embodiment

The present invention provides an apparatus that is useful in electrochemical cells, especially those useful for generating electricity known as fuel cells. More particularly, the invention provides various electrode structures or configurations that provide increased power output per unit weight of the cell. The electrodes may be used with gaseous fuels, such as hydrogen, or liquid fuels, such as methanol. The electrodes are designed to operate even at low fuel and oxidant pressures.

One aspect of the invention provides an electrically conductive member disposed within the electrode, rather than having a conductive cloth layer and a

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conductive frame disposed along a perimeter face of the electrode The electrically conductive member may be a sheet of expanded metal mesh or wire, preferably expanded metal mesh. The expanded metal mesh or other electrically conductive member preferably has a greater portion of open area than does carbon cloth or paper to increase the gas flow to and from the catalyst areas.

Another aspect of the invention provides a wicking member for delivering a liquid fuel to the anode side of the fuel cell or to the electrochemical cell. The liquid fuel may be mixed with various concentrations of water as necessary to facilitate the electrode reactions and provide water to the proton exchange membranes.

A further aspect of the invention provides a cooling member disposed between adjacent cells. While cooling may be accomplished by external air circulation, either natural or forced, this is sometimes inadequate, and it is sometimes advantageous to add a cooling member. The preferred cooling members of the invention comprise air channels disposed between fuel feed ports in the anode compartment. The channels may be as simple as a pair of walls with an opening therebetween or the channel may contain one or more heat transfer members.

The present invention provides a lightweight fuel cell system that operates with fuel and oxidant gases at near atmospheric pressures. The fuel cell system uses a monopolar cell design where the electrode surfaces are sufficiently accessible to the gases that it is not necessary for the oxidizer and reducer gases to be compressed. The fuel cell is ideal for personal use due to its light weight, compact size and self contained operation. An added feature of the present invention is that individual fuel cells may be linked together to achieve greater voltages or currents.

The present invention provides a monopolar fuel cell system based on the use of flat cell frames that support the cell components. A monopolar design is lighter than a bipolar or filter press design for a number of reasons. First, since the monopolar fuel cell is not under compression, the components, such as the cell frames, can be much thinner than the structure of a bipolar fuel cell which relies on the thickness of the components to maintain precise spacings for gas flow therethrough. The primary limitation on the thickness of the components is that the component in question must have sufficient strength to maintain its integrity during use in various

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applications. This permits the use of a wide variety of lightweight conductive materials, such as a thin titanium sheet.

Another reason that the monopolar fuel cell of the present invention weighs less than conventional systems, is that many components of a conventional system are no longer necessary and may be eliminated entirely. For example, the preferred monopolar fuel cell uses ambient air as the oxidizer, rather than compressed air or oxygen, as the oxidizer. Because it is unnecessary to compress the oxidizer, the conventional gas control system is unnecessary and may be eliminated. Instead, the gas is supplied to the cathode by either free or forced convection. In either case, there is no need for an enclosed cathode flow field.

In addition, the bipolar plates of conventional fuel cells, which separate anode and cathode flow fields and conduct current from one cell to the next, are no longer needed and may be eliminated. The plates are not needed to separate flow fields because the anode and cathode flow fields of the present invention are open to the sealed reducing gas chamber and the atmosphere, respectively, and are effectively separated by the cells themselves. Furthermore, the bipolar plates are not needed to conduct electrical current from one cell to the next since this function is performed by a different electrically conducting member, such as an external bipolar frame or an internal bipolar grid.

Another component or system of conventional fuel cell systems that is no longer required is the heat transfer system. The fuel cell system of this invention has a much larger ratio of exposed area to active area and, therefore, dissipates waste heat directly to the surrounding air without the need for a circulating heat transfer fluid, a pump to circulate the fluid, or heat exchangers inside the stack to collect the heat and outside the stack to dissipate the heat. However, in some applications, it may still be desirable to include a cooling member such as an air channel made from a material having a high thermal conductivity.

Eliminating moving parts from fuel gas systems, in accordance with the present invention, eliminates the parasitic power losses required to drive the motors involved and the potential for malfunctions. Furthermore, eliminating small motors and moving parts in fuel gas systems reduces maintenance and malfunctions, particularly in

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applications where the system is portable.

Another advantage of monopolar fuel cell systems is the low cost of manufacture. Because the entire stack is assembled from a series of flat, or nearly flat, components, all of the parts can be cut from flat sheets. No complex machining steps are required.

Electrochemical Cells Having External Bipolar Frames

The membrane and electrode assemblies used in the present invention are comprised of a proton exchange membrane (PEM), catalyst layers on either side of the PEM, and gas diffusion layers over the catalyst layers. Gas diffusion electrodes, their construction and fabrication, are generally described in Murphy et al., U.S. Patent No. 5,460,705 which is hereby incorporated by reference. The preferred PEM is generally a polymer material having sulfonate functional groups contained on a fluorinated carbon backbone, such as the perfluorinated sulfonic acid polymers available under the trade name NAFION from Du Pont de Nemours, E. I. & Co of Wilmington, Delaware. The gas diffusion layers preferably comprise a carbon conducting paper or cloth. In one preferred embodiment, a platinum catalyst is supported on the carbon paper before hot pressing with the PEM. However, it is also anticipated that the same general construction can be achieved using thin film electrodes, where the catalyst is deposited directly onto the PEM rather than the carbon paper.

It is preferred that the reducing gas be delivered to the anodes from a supply vessel through a pressure regulating feed valve and feed line connected to the reducing gas chamber. The anode surface of each individual cell is directly exposed to the reducing gas in the chamber and does not require enclosed anode flow fields for delivery of the reducing gas. Since the reducing gas chamber allows the reducing gas to flow freely over the anode surface, it is not necessary to compress the reducing gas for delivery through narrow passages or flow fields. Therefore, the reducing gas may be supplied at any pressure. However, it should be noted that while the fuel cell does not require the reducing gas, such as hydrogen, to be pressurized, it is generally preferred that the hydrogen is stored in a pressurized vessel that can be transported along with the fuel cell itself. It is further preferred that the pressurized hydrogen be

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delivered from the vessel to the reducing gas chamber through a step-down regulator to a pressure generally below about one atmosphere, but most preferably below about 2 psi to avoid displacing the membrane and electrode assemblies out of their frames.

The fuel cell consumes the fuel at the anode, produces water at the cathode and generates a flow of electricity for use in various applications. The water that is 5 generated at the cathode is useful to keep the PEM moist so that it will conduct protons efficiently. Water will evaporate from the cathode surface into the air and provide some cooling to the cells. However, because there is no external source of water to the PEM, the air flow rate and temperature should not be allowed to dry out the PEM

In one aspect of the invention, it is possible to fabricate the cells with the same catalyst, preferably platinum (Pt) or a platinum-containing alloy, and the same catalyst loading on both the anode and the cathode. Therefore, the cells may be cut, handled and fabricated without regard for their orientation. This may be advantageous in some applications, since the anode and cathode cannot be distinguished by appearance alone. On the other hand, because Pt has greater catalytic activity for hydrogen gas than for oxygen gas, a cell with better precious metal utilization is obtained when the platinum loading is lower on the anode than on the cathode.

The sheets and other components of the present invention may be assembled and coupled in many different ways and with many different means as will be recognized in the art. In particular, it should be recognized that the components may be fastened with mechanical fasteners, such as bolts, screws, clamps, straps, rivets and the like, adhesives, and welding of the components. However, it is most preferred that all of the components be fastened using adhesives and welding, so that a minimal amount of material is added to the fuel cell and a minimal number of parts are needed. Bonding processes are also generally preferred since this type of assembly can easily be automated

Now referring to Figure 3, a face view of three types of cell frame components are shown. First, there is the dual cell frame 30 which has openings 32, 34 for exposing two M&E assemblies. The dual cell frames 30 are overlapped (as shown in Figure 4) so that the opening 32 of a first frame 30 coincides with the opening 43 of an

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adjacent frame 30. The two metal frames 30 hold the M&E assemblies firmly around the edge of the assembly.

Now referring to Figure 4, a top cross-sectional view of a fuel cell stack 50 is shown. The overlapping metal frames 30 can be repeated any number of times to form the multi-cell stack or fin 51. At the end of the series of cells, a terminal cell frame 36 may be used to overlap one of the openings 32, 34 and also to provide an electrical contact 37 for conducting electricity to and from the stack. The curved dual cell frame 38 has two openings 40, 42 like the dual cell frame 30, except that it has an enlarged central region 44 which can be curved to form the distal end of a fin 51. The fin 51 can have any number of cells 52, but is shown here having eight cells connected in series. In a fashion similar to the M&E assembly of Figure 2, the cells 52 have an M&E assembly comprised of a central PEM 54, a catalyst coating 56 and gas diffusion layer 58.

Side plates 60 are sealed to the top and bottom edges of the fin 51 and an end cap 62 is sealed to the open end of the fin 51. The plates 60 and cap 62, in cooperation with the fin 51, form a chamber 64 in communication with all of the anode surfaces 66. A gas inlet port 68 is provided in the end cap 62 for the introduction of a fuel to the chamber 64.

The two-sided arrangement shown in Figure 4 represents a preferred embodiment of the invention. This arrangement permits the largest number of cells to share a common internal fuel gas chamber. It is also possible to arrange the series of cells as a flat strip, where the opposing surface of the internal chamber is an inactive surface. While generally less efficient in terms of weight or volume, in some applications, this may be a preferable arrangement.

The preferred fuel is a reducing gas, with the most preferred gas being hydrogen. In operation, each of the individual cells of the fuel cell 50 has a cathode surface 70 that is exposed to the air as the oxidizing gas. Therefore, the fuel cell 50 operates at low pressures and without any moving parts. The cell is also lightweight, easy to manufacture and relatively inexpensive.

Unlike a conventional bipolar system, this fuel cell collects the current only from the edges of the electrode, leaving the entire face open. Leaving the area of the

electrode as open as possible reduces the potential for interference with diffusion and facilitates the supply of air under ambient conditions without the need for a compressor or other high powered air moving device.

A four cell version of fuel cell 50 of the present invention was constructed and operated using hydrogen gas as the fuel and ambient air as the oxidizer. The performance of the fuel cell is illustrated in Figure 5 by a graph of stack potential (Volts) as a function of stack current (Amps). As is apparent from the data displayed in that figure, increasing the fuel pressure slightly (1.5 psig) and using a small fan to create a forced convection flow over the stack produced a marked improvement in the stacks performance.

Now referring to Figures 6(a-c), the structures of three different M&E assemblies specifically intended for use in a monopolar fuel cell are shown. In the configuration of Figure 6(a), a metal grid 82 is embedded in the front surface of the electrode 80 and in direct contact with both the electrocatalytically active portion 56 of the electrode and the membrane 54. The advantage of this arrangement is good electrical contact with the electrocatalyst for efficient current collection and the least interference with gas diffusion within the electrode. The disadvantages are the difficulty in making good electrical contact between the current collection plates 30 and both the metal grid 82 and the conductive carbon cloth 84 at the back of the electrode, the potential for damaging the membrane 54 with the grid while attaching the electrocatalyst and the membrane. Although the latter can be partly compensated for by appropriate impregnation of the electrode with a NAFION solution, the long diffusion path for protons around the grid through the recast membrane leads to relatively high local resistances.

Figure 6(b) shows an M&E configuration 85 with the metal grid 82 in the center of the electrode, between the uncatalyzed carbon gas diffusion layer 58 and the active electrocatalyst layer 56. The advantages of this configuration are good contact between the grid 82 and the electrocatalyst 56 for efficient current collection and no blockage of the interface between the membrane 54 and the active electrocatalyst 56. The disadvantages of this configuration are a more difficult fabrication procedure and

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possible interference with gas diffusion inside the electrode.

Figure 6(c) shows an M&E configuration 90 with the grid 82 on the back side of the electrode. The advantages of this configuration are good contact between the grid 82 and the cell frame 30 (see Figure 4) for efficient current collection, no blockage of the interface between the membrane and the active electrocatalyst, and simpler fabrication than the arrangement with the grid inside the electrode. The disadvantages of this arrangement are the creation of vertices on the outer face of the electrode which can act as anchors for water drops leading to a loss in available area for gas diffusion and the difficulty of insuring that the grid will stay in contact with the electrode for an indefinite period of time.

Electrochemical Cells With Internal Bipolar Electrically Conducting Member

One aspect of the invention provides an electrically conductive member disposed within the electrode, rather than having a conductive cloth layer and a conductive frame disposed along a perimeter face of the electrode. The electrically conductive member may be a sheet of expanded metal mesh or wire, preferably expanded metal mesh. The expanded metal mesh or other electrically conductive member preferably has a greater portion of open area than does carbon cloth or paper to increase the gas flow to and from the catalyst areas.

Now referring to Figures 30(a-b), the structures of two different M&E assemblies specifically intended for use in a stack of monopolar fuel cells wherein the internal electrically conductive element 82 of the anode in one cell extends and forms the conducting element in the cathode of an adjacent cell.

The difference between the M&E configurations of Figures 30(a-b) and those of Figures 6(a-b) respectively is the elimination of the conductive carbon cloth 84 shown at the back of the electrodes in Figures 6(a-b).

The metal grids of Figures 30(a-b) provide the support normally provided by the carbon cloth or carbon paper in conventional gas diffusion electrodes while adding in-plane conductivity to the electrode. Since the metal grid is quite close to the membrane in this design, it is imperative that the metal not corrode. Corrosion will not only increase the contact resistance between the active portion of the electrode and the

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current collecting frame, but mobile metal ions coming in contact with the membrane may replace protons in the membrane. Replacing even a small fraction of the protons in the membrane with far less mobile metal ions will lead to a significant drop in membrane conductivity. This factor imposes a requirement that the grids have low contact resistance and be corrosion resistant. The best way to impart these properties to a piece of lightweight material is to plate the metal with a layer of a more precious metal, such as gold, platinum, palladium, or ruthenium, to protect it from corrosion and to improve electrical contacts.

Another advantage to fabricating electrodes with an internal conductive element so long as the conductivity of the resultant electrode is sufficient, is that the bipolar cell frame described above is no longer needed for consecutively connecting the cells. Eliminating the cell frame leads to a smaller and lighter stack.

Figure 7 shows the improvement in monopolar fuel cell performance produced by the inclusion of a metal grid on the back of the electrode. The configuration 90 of Figure 6(c), with the grid on the back side of the electrode, gave the best performance of the three designs in Figures 6(a-c).

Figure 8 shows a module design 95 that would stack head-to-tail (end 96 of one module to end 98 of another module) using a combined plug 99 with four electrical connections 94 and a gas connection 99. Inside the module, terminal ends of the cells are electrically connected into groups, each of which will supply power at a potential of 12 Volts (the minimum output that a power system may have for many applications). The mating plug on the item being powered (not shown) can be wired to utilize the groups in parallel for 12 Volts, or in series for 24 Volts. As many modules 95 as desired can be connected together to increase the current available, but the voltage will remain constant at either 12 or 24 Volts. It is anticipated that the modules could be stacked side-by-side or end-to-end and operate equally well.

The present invention provides an open fin design that produces a very light fuel cell system. The power-to-weight ratio for the fins alone (no fittings) has been shown to be as high as 220 W/lb. When the weight of the fittings needed to connect the stack to the fuel supply and the next module are included, this drops to a still impressive 180 W/lb. While this is the lightest arrangement, this design is only

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practical for enclosed applications because the lightweight construction leaves the cell subject to damage.

Figure 9 shows an even more compact and useful fuel cell configuration 100. In this arrangement a plurality of fins 102 are enclosed in a common housing or shell 104, with a small fan 106 located at one end to move a continuous stream of air over the cathodes 108. With 4 mm thick fins, a fin-to-fin spacing of 2-3 mm would be required, or about 4 fins per inch. This arrangement produces a projected stack volume of about 335 ml for a module sized to produce about 50 Watts of power. It is likely that this volume can be reduced further, with a volume below about 300 ml possible.

The fuel is delivered to each anode chamber 110 through a manifold 112, preferably located at the rear of the fins. A fuel source, such as a pressurized hydrogen tank and step-down pressure regulator (not shown), is connected to the coupling 114 on the manifold. With hydrogen in communication with the anodes and air passing over the cathodes 108 and out the back grill 116 of the housing 104, the fuel cell 100 will generate electricity for use in any 12 or 24 Volt DC application.

The addition of the fan 106 and the shell 104 increase the weight of the system 100 slightly, thereby reducing the power/weight ratio to about 150 W/lb for a 50 Watt module. Although the fuel cell 100 represents an increase in the amount of weight needed for any specified power output, the fuel cell is still quite light and the smaller size makes the cell much more portable and easy to carry.

The light weight fuel cell design of the present invention is suitable for use with gaseous fuels, such as hydrogen, or with liquid fuels, such as methanol, dimethoxymethane. or trimethoxymethane.

Both of these systems, as shown in Figures 8 and 9, are drawn as hydrogen fuel cells, with the gas being supplied through a feed line 94. This technology can be used with liquid fuels as well. In the case of liquid fuels, the water-alcohol mixture consumed by the fuel cell would be supplied using the same common manifold fuel distribution approach as shown for hydrogen. Attitude independence, easily achieved in a gas fueled system, can be achieved in a liquid fueled system through the use of a

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wicking system to feed the anodes by using capillary action to lift the fluid from the bottom of a liquid fuel container (not shown), where ever it may reside.

Figure 10 is an exploded view of a subassembly 120 including a pair of gas diffusion electrodes 122 fabricated with internal bipolar conductive metal grids, made in accordance with Figure 6(b). The electrodes 122 each comprise an active cathodic catalyst region 126, an active anodic catalyst region 128 and a gas barrier 130 disposed therebetween. A proton exchange membrane 132, preferably having a PTFE frame 134 is disposed between the cathodic region of one electrode and the anodic region of another electrode. While each active catalytic region is shown as having the catalyst deposited on one surface of the gas diffusion matrix, it is also possible to deposit a thin layer electrode on the surface of the membrane. In the former situation, one surface of the gas diffusion electrode is catalyzed on one side of the gas barrier and the opposite surface is catalyzed on the other side. In the latter case, the diffusion backing is bonded to the back of the electrode, and the gas diffusion matrix can be fabricated as a symmetrical unit, i.e., it is not necessary to deposit catalyst on either side thereof

In either case, the electrodes are assembled with a PEM membrane between them and a PTFE "window frame" gasket included to eliminate the possibility of electrodes shorting at the edge due to physical contact of the anode 128 and cathode 126. In this geometry each grid has both an anode and a cathode bonded to it and serves as the bipolar link between them. This eliminates the need for the bipolar cell frames. In this design the conductive elements within the electrodes serve as the bipolar elements connecting each pair of adjacent cells. PTFE frames are included to prevent shorting from electrode to electrode at the edge.

The gas barrier strip 130 disposed down the center of the conductive metal grid can be fabricated in a variety of ways. A polymer strip can be deposited using a self curing polymer, such as silicones, epoxies, and urethanes; a thermoplastic; or easily melted metals, such as solder. The barrier can also be produced as part of the grid fabrication process with the gas barrier strip 130 comprising a region of unexpanded metal. It can be readily appreciated by those skilled in the art that other

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methods may be used in producing light weight fuel cells, such methods are considered to be within the scope of the present invention.

Also, a wide variety of materials may be suitable for use as the conductive metal element within the gas diffusion electrode. Those described here are only examples, the use of materials not described here is within the scope of the present invention. The preferred material for the conductive element is expanded metal, a product fabricated by piercing and stretching a sheet of metal or metallic foil. For optimal performance, the metal should be flattened after expansion to restore it to its original thickness.

Figure 11 shows the basic form of the expanded metal sheet used in conjunction with this invention. Expanded metal foils with the pattern shown as well as others are available from a wide variety of manufacturers, in a wide range of thickness, from 0.003" (0.076 mm) to 0.1" (2.5 mm) or more and manufactured from a variety of metals, including, but not limited to, titanium, nickel, copper, stainless steel, aluminum, and niobium. They are available with open areas (the percentage of the sheet area comprised of the holes) ranging from 10% to 70%. A typical expanded metal sheet has two primary directions. For best performance, the expanded metal sheet should be oriented such that the current flows in the direction parallel to the "long way" of the diamond, since the electrical resistance is lowest in this direction, as shown in Table I, which contains some properties of typical expanded metals and other materials useful for forming the conductive portion of the electrode structures, as taught here. Some of the materials in this table were gold plated prior electrical measurements so that contact resistance is reduced.

Table I. Properties of Metal Conductive Components

Label	Thickness	LWD*	SWD*	Areal	Resistivity	Resistivity	Open
	(cm)	(cm)	(cm)	density	(LWD)	(SWD)	Fraction
				(g/cm ²)	(mΩ cm/cm)	(mΩ cm/cm)	(%)
Ni #1	0.028	0.167	0.125	0.01994	6.08	9.62	51
Ni #2	0.028	0.172	0.125	0.01977	5.72	10.34	56.5
Ni #3	0.0285	0.325	0.112	0.03046	. 2.79	10,0	43

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0.016	0.186	0.110	0.0493	1.78	4.68	35
0.041	0.185	0.113	0.0383	2.62	9.25	41
0.009	0.138	0.075	0.0093	35	119	52
0.014	0.123	0.071	0.0288	. 33	82	31.4
0.009	0.1	0.079	0.0185	5 0	133	34
0.011	0.134	0.1	0.0219	42	123	40
0.0172	0.14	0.0074	0.0545	0.38	1.17	31
0.017	0.2	0.09	0.06958	0.26	0.93	37.8
0.05	n.a.°	n.a.c	0.0853	3.2 ^d	n.a.c	68 ^e
0.0135	n.a.°	n,a.°	0.06	6.7 ^d	n.a.c	_38 ^f
	0.041 0.009 0.014 0.009 0.011 0.0172 0.017 0.05	0.041 0.185 0.009 0.138 0.014 0.123 0.009 0.1 0.011 0.134 0.0172 0.14 0.017 0.2 0.05 n.a.s	0.041 0.185 0.113 0.009 0.138 0.075 0.014 0.123 0.071 0.009 0.1 0.079 0.011 0.134 0.1 0.0172 0.14 0.0074 0.017 0.2 0.09 0.05 n.a.c. n.a.c.	0.041 0.185 0.113 0.0383 0.009 0.138 0.075 0.0093 0.014 0.123 0.071 0.0288 0.009 0.1 0.079 0.0185 0.011 0.134 0.1 0.0219 0.0172 0.14 0.0074 0.0545 0.017 0.2 0.09 0.06958 0.05 n.a* n.a* 0.0853	0.041 0.185 0.113 0.0383 2.62 0.009 0.138 0.075 0.0093 35 0.014 0.123 0.071 0.0288 33 0.009 0.1 0.079 0.0185 50 0.011 0.134 0.1 0.0219 42 0.0172 0.14 0.0074 0.0545 0.38 0.017 0.2 0.09 0.06958 0.26 0.05 n.a.c. 0.0853 3.24	0.041 0.185 0.113 0.0383 2.62 9.25 0.009 0.138 0.075 0.0093 35 119 0.014 0.123 0.071 0.0288 33 82 0.009 0.1 0.079 0.0185 50 133 0.011 0.134 0.1 0.0219 42 123 0.0172 0.14 0.0074 0.0545 0.38 1.17 0.017 0.2 0.09 0.06958 0.26 0.93 0.05 n.a* n.a* 0.0853 3.2* n.a*

Notes:

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- a LWD and SWD are defined in Figure 11.
- b Not gold plated.
- Not applicable, this material is not asymmetric.
- d Resistivity in either direction, material is symmetric.
 - e Calculated
 - f Manufacturer's value.

Perforated metal sheets are also suitable as the conductive component within the electrode. Compared to expanded metal, perforated metal sheets are generally stronger and more conductive, but generally display a smaller open fraction (<40%) and are thus less conductive to gas exchange.

Woven metal wire products are suitable as well. Compared to expanded metal, woven products have more open area (up to 80%) for superior gas exchange. However, the weaving process makes woven metal wire products more expensive to manufacture than expanded metal, and, because electrical conductivity requires that the current flow through a large number of wire-to-wire contacts, resistivity is high. Furthermore, because of the woven nature of the material, any individual sheet can only be as thin as twice the wire diameter.

20 Conventional carbon cloth supported gas diffusion electrodes have a gas diffusion matrix consisting of conductive carbon powder bound together, and to the carbon cloth, by polytetrafluoroethylene (PTFE). Since the expanded metal has substantially more open area than the carbon cloth, a modification of the gas diffusion matrix formulation would be advantageous. It has been found that replacing a portion of the carbon powder with finely cut and divided conductive carbon fibers produces a superior gas diffusion matrix. The inclusion of conductive carbon fibers in the gas diffusion matrix improves the matrix's ability to span the large gaps in the expanded metal, or other open metal, used as the conductive support.

Figures 12(a-c) are three views of a fin configured for liquid fuel use. Figure 12(a) is a top view of a liquid fuel cell system 100 showing the top view of a fin 101 defining a cathodic surface 103. Figure 12(b) is a cross-sectional view of the fin 101 cut vertically near the center of the arrangement shown in Figure 12(a). The anodes 104 in the fin 101 define anodic surfaces 105 that are provided with liquid fuel. Figure 12(c) is an enlargement of a portion cut out of the fin 101 as presented in Figure 12(b). The wick structure 106 is designed so that all of the anodes 104 are supplied with fuel at all times. Because of the high energy storage density associated with liquid fuels, the volume inside the fin allows the storage of sufficient fuel to operate for an extended period of time, from several hours to a day, depending on the load. This eliminates the need for any other fuel storage in such a system, unless an even more extended operating period is required. Refueling the liquid fueled system is equally easy, requiring only a container of a nourable liquid

Fuel cell designs in accordance with the present invention are useful in a wide variety of configurations.

Figures 31(a-c) are three views of a fin depicting a compact fuel cell arrangement. In this arrangement, two sets of cells, fabricated using a common conductive element in the electrodes as the polar element, are attached to opposing faces of a single polymeric frame. configured for liquid fuel use.

Figure 31(a) is a top view of a hydrogen/air fuel cell system 310 showing the top view of a fin 311 defining a cathodic surface 313.

Figure 31(b) is a cross-sectional view of the fin 311 cut vertically near the center of the arrangement shown in Figure 31(a).

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Figure 3 (c) is a cross-sectional view cut horizontally near the center of Figure 31(a).

Figure 25 is a schematic diagram depicting a polymeric frame 230 that can be used in conjunction with the present invention. The polar frame comprises a first passage 232 for hydrogen feed, a central element 234, and a second passage 236 for hydrogen exit. The central element serves both in preventing contact between the electrodes and channeling the hydrogen in a U-shaped course through a crossover channel 238. It can be easily appreciated that non-polymeric frames can also be used with the present invention. However, when conducting frames are used, they must be insulated to prevent the formation of a short circuit when the frame contacts the face of the anodes.

In the arrangement schematically depicted in Figure 31, a large fraction of the external surface of the fuel cell assembly serves as active cathode area. The fins of Figure 31 can be used as the basic structural element in larger arrangements such as the ones depicted in Figures 8 and 9. These fins can also be operated individually in small power supplies. While the fin of Figure 31 comprises two sets of 5 cells each for a total number of 10 cells connected in series, there are no limitations on the number of cells that can be connected in the manner described here. Arrangements with a larger or a smaller number of cells are considered within the scope of the present invention. Also, It is not a necessary element of this invention to make symmetrical fins with the same number of cells on two sides, non-symmetrical arrangements are variations within the scope of the present invention. The arrangement described in this invention presents high flexibility. Adjustments in the number of cells and their arrangement are provided for by the present invention so that optimum use is obtained in any particular application.

The significant advantages provided by the compactness in the arrangement presented in this invention may be offset by the difficulty in dissipating the heat generated by operating such compact systems. Another aspect of the invention is to provide means for avoiding the loss in efficiency that may occur when the fuel cell system is operated for longer periods of time.

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Figure 26(a) is a cross sectional view of a segment of the fin of Figure 31(b) comprising two membrane and electrode assemblies 244 positioned opposite each other with an internal heat transfer structure 242 positioned between the hydrogen feeds 240. The polymer frames 246 are also shown. Figure 26(b) is an edge view of the fin of Figure 12(b) showing the internal heat structure 242.

Figure 27 is a drawing depicting a wavy convoluted aluminum sheet 250 as one example of internal heat transfer structures that can be used in conjunction with the present invention. The sheet provides one example for the inclusion of a cooling channel and a heat transfer element in the fin of Figure 12. In this instance, the sheet is preferably oriented so that the flow direction of the channels in the sheet are parallel to the longest dimension of the fin. Those skilled in the art of heat transfer can easily appreciate that other orientations of the sheet or the use of an element made of a different metal or presenting a different pattern provide internal heat transfer structures that can be used with the cooling system described here. The presence of an internal heat transfer structure is not a necessary element of the cooling system disclosed in this invention. Effective cooling may be obtained even if the fin is fabricated with a channel that has no heat transfer element.

While the examples discussed below describe cooling systems that present a cooling flow parallel to the long direction of the fin, equal cooling efficiency is obtained when the cooling flow is parallel to the short direction of the fin. One practical advantage presented by arrangements having parallel flow of cooling air and reaction air is the possibility to use one fan for producing air flow that can be used for both purposes. The advantage in using a single fan can be offset when independent control of the cooling and reaction flows is desirable. On the other hand, fuel systems with cooling and reaction flows that are perpendicular present the advantage of independently controlling the flows but require at least two air moving devices. The choice between an arrangement with parallel cooling and reaction flows and an arrangement with perpendicular flows depends on the particular application.

Another aspect of the invention is to provide for protection of the exposed cathodes from damage by foreign objects. Such protection can be achieved by adding a protective screen to the arrangement described in Figure 26. Figure 28 is a cross

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sectional view of the fin of Figure 26(a) with the back plane of the monopolar cells extended and folded over to form a protective cover. The fin comprises M&E's 264, hydrogen manifolds 266, a cooling channel with a heat transfer element 268, and ventilated protective covers 262. In one embodiment of the invention, the protective cover is formed by extending the metal sheet serving as the backside of the hydrogen manifold, perforating it, and bending it around to form a cover over the faces of the cathodes. By using only a single sheet of metal as both the backside of the manifold and as the protective structure, thermal conductivity is maximized. An additional advantage of the present design is the contribution of the protective structure to heat dissipation during the operation of the fuel cell system. The portion of the fin serving as a protective cover comprises an open area so that efficient air exchange with the atmosphere is maintained. The open area may be obtained by placing perforations in the portion of the fin that serves as a protective cover. Other means for maintaining gas exchange between the cathodes and the atmosphere are within the scope of the present invention.

Figure 29 is a cross sectional view of a fin with protective cooling fins. The cathodes 274 are protected by protecting cooling fins 272 that are bent and form bumpers at the edge of cooling channel 278. In this arrangement, the bumpers offer protection to the cathodes while serving as the heat transfer element. The arrangement also presents the advantage of providing a fuel cell system with cathodes that have maximum access to the atmosphere.

Other arrangements included within the scope of the present invention are shown in Figures 13(a-b). Figures 13(a) and 13(b) are side views of two configurations in which the cells are wrapped around the fuel tank 118. Figure 14 is a cross-sectional view of a monopolar fuel cell system 120. Curved monopolar fuel cell frames 122 and cured membrane and electrode assemblies 124 are mounted on the outside of a vessel 128 that serves as a fuel reservoir for storing the fuel to be consumed by operating the fuel cell system. When the fuel cylinder is a storage vessel filled with liquid fuel, the operating time for this configuration would be much longer than when hydrogen gas is used as the fuel. It is important to note that in this invention, the conceptual difference in the design of gas and liquid fueled cells is the

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addition in liquid fuel cell designs of a wicking element inside the open volume reserved for hydrogen delivery in the gas fueled cell design.

Given the high energy density of liquid fuels, a monopolar fuel cell of the design taught here can be fabricated in a wide variety of shapes that can be integrated into the device that they are intended to power. All of the needed fuel can be stored within the fuel cell structure (i.e., no external tank). For example, a methanol-fueled, 5 Watt flat panel fuel cell with a 12 cell array, operating at a potential of 0.5 V/cell and a current density of 60 mA/cm² occupies an area of about 100 cm² and is 7.5 mm thick. The fuel stored inside such a cell is adequate for over 20 hours of operation

The examples shown here are all illustrative of fuel cells, but the present invention can be useful in other applications as well. For example, by filling the wick shown in Figure 12 with water, the stack can be operated as an electrolyzer to generate hydrogen at the cathode and oxygen at the anode. It can be readily appreciated that various other applications of the concepts described here are within the scope of the present invention.

The following examples show the function of this invention and some of its preferred embodiments.

Example 1

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This example illustrates a gas diffusion electrode using carbon powder as the conductive material in the metal supported matrix.

A gas diffusion electrode was made by mixing Vulcan XC-72R high surface area carbon powder (available from CABOT Technology Division, Pampa, Texas) with polytetrafluoroethylene "PTFE" suspension (such as T-30, available from DuPont, Wilmington Delaware) in a 60:40 ratio (based on the dry weight of the PTFE), water, and a nonionic surfactant (such as Triton X100, available from Fisher Scientific, Fair Lawn, New Jersey). The carbon/PTFE mixture was sonicated to reach complete dispersion and the resulting paste was spread onto an expanded foil grid. This grid, fabricated from expanded copper foil, had previously been electroplated with a protective gold layer, through the use of a standard commercial gold

electroplating both (E-56, from Engelhard, Corp., Iselin, N.J.) and serves as the grid to form a gas diffusion electrode.

The bulk of the water was removed from the gas diffusion electrode by filtration. Polycarbonate filter membranes (5 µm pore size, available from Poretics, Livermore, California) were found to give the most efficient water removal, with only traces of carbon occasionally passing through and easy separation of the electrode from the filter. Finer pore materials (1 and 3 µm) blocked all of the carbon, but had much lower rates for water removal. Other filter materials did not release well from the electrode and retained carbon on the surface of the filter. The use of a heat lamp to gently warm the electrode during the filtration step was found to improve the water removal. The electrode was dried at room temperature in a vacuum oven oven for 4 to 5 hours, then further dried at 60 °C in the same vacuum oven overnight. Finally, the electrode was treated in an argon atmosphere at 320 °C for 2 hours in order to sinter the PTFE and decompose the surfactant. The density and thickness of the final gas diffusion electrodes were about 0.57 g/cm³ and 0.7 mm, respectively.

Example 2

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This example illustrates a gas diffusion electrode using carbon fiber with carbon powder in the metal supported matrix.

The gas diffusion electrode was made by mixing Vulcan XC-72R high surface area carbon powder (CABOT) and Thornel™ DKD-X carbon fiber (Amoco Corp., Apharetta, GA) with PTFE suspension (T-30, DuPont), in a 52:17:31 ratio (based on the dry weight of the PTFE), water, and nonionic surfactant (Triton X100). The carbon/PTFE mixture was sonicated and fabricated into a metal supported gas diffusion electrode as described in example 1.

Example 3

This example illustrates a gas diffusion electrode using carbon fiber with carbon powder in the metal supported matrix.

The gas diffusion electrode was made by mixing Vulcan XC-72R high surface area carbon powder (CABOT) and carbon fiber obtained by ball milling Toray TGP-H

paper (Toray Corp., New York, NY) with PTFE suspension (T-30, DuPont), in a 42.17:41 ratio (based on the dry weight of the PTFE), water, and nonionic surfactant (Triton X100). The carbon/PTFE mixture was sonicated and fabricated into a metal supported gas diffusion electrode as described in example 1.

Example 4

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This example illustrates a gas diffusion electrode using carbon fiber with carbon powder in the metal supported matrix.

The gas diffusion electrode was made by mixing Vulcan XC-72R high surface area carbon powder (CABOT) and Thornel™ DKD-X carbon fiber (Amoco Corp.) with PTFE suspension (T-30, DuPont), in a 33:16:51 ratio (based on the dry weight of the PTFE), water, and nonionic surfactant (Triton X100). The carbon/PTFE mixture was sonicated and fabricated into a metal supported gas diffusion electrode as described in example 1.

Example 5

This example illustrates a gas diffusion electrode using carbon fiber with carbon powder in the metal supported matrix.

The gas diffusion electrode was made by mixing Vulcan XC-72R high surface area carbon powder (CABOT) and Thornel™ DKD-X carbon fiber (Amoco Corp.) with PTFE suspension (T-30, DuPont), in a 40:20:40 ratio (based on the dry weight of the PTFE), water, and nonionic surfactant (Triton X100). The carbon/PTFE mixture was sonicated and fabricated into a metal supported gas diffusion electrode as described in example 1.

The compositions from the first five examples are summarized in Table II.

Table II. The Compositions Of The Gas Diffusion Electrode in Examples 1

Example	XC-72R carbon	Thornel carbon	Ground		Triton	
	powder	fibers	Toray paper	PTFE	X100	
1	60%			40%	2% of C	

2	52%	17%		31%	2% of C
3	42%	1	17%	41%	2% of C
4	33%	16%		51%	2% of C
5	40%	20%	1	40%	2% of C

Example 6

This example demonstrates an alternative method for forming the gas diffusion matrix

The gas diffusion matrix can also be fabricated by a direct filtration approach where the metallic conductor is placed on a polycarbonate filter and the gas diffusion matrix impregnated into the conductor by filtration from a more dilute suspension. Vulcan XC-72R high surface area carbon powder, conductive carbon fiber (ThornelTM DKD-X), and PTFE suspension, (T-30), were mixed in a 33:16:51 ratio (based on the dry weight of the PTFE) with water and nonionic surfactant. The mixture is sonicated to separate and disperse all of the particles into a uniform suspension. This mixture is then filtered through a cleaned expanded nickel conductive element placed on a polycarbonate filter membrane (1 µm pore size) to remove water and directly form a gas diffusion layer around the metallic conductor.

Example 7

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This example demonstrates a method for forming the active portion of the electrode.

The thin film catalyst layer for the electrode was made by mixing platinum black catalyst (such as fuel cell platinum black catalyst available from Engelhard, Iselin, New Jersey) with a 5 % solution of 950 equivalent weight Nafion[®] (available from Solution Technology, Inc., Mendenhall Pennsylvania). Sufficient water was added to produce a viscous, but fluid mixture. The mixture was then sonicated to disperse the platinum black and give a uniform ink. This ink was brushed directly onto the Nafion[®] membrane and dried.

Example 8

This example demonstrates an alternative method for forming the active portion of the electrode.

The thin film catalyst layer for the electrode was made by mixing platinum black catalyst (Engelhard's fuel cell grade) with a 5 % solution of 950 equivalent weight Nafion[®] (available from Solution Technology, Inc., Mendenhall Pennsylvania). Sufficient water was added to produce a viscous, but fluid mixture. The mixture was then sonicated to disperse the Platinum black and give a uniform ink. This ink was brushed directly onto a gas diffusion matrix which had been produced as described in Example 7 and dried at under ambient conditions.

Example 9

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This example demonstrates still another method for forming the active portion of the electrode.

The catalyst layer for the electrode was made by mixing platinum black catalyst (Engelhard's fuel cell grade) with a PTFE suspension (T-30). Sufficient water was added to produce a viscous, but fluid mixture. The mixture was then sonicated to disperse the platinum black and give a uniform ink. This ink was brushed directly onto a gas diffusion matrix which had been produced as described in Example 7. The electrode was allowed to dry under ambient conditions until it appeared dry to the eye, then further dried at 60 °C in a vacuum. After vacuum drying the electrode was heated to 320 °C in a flowing argon atmosphere and held at that temperature for 10 minutes to sinter the PTFE and decompose the surfactant

25 Example 10

This example demonstrates still another method for forming the active portion of the electrode.

The thin film catalyst layer for the electrode was made by mixing platinum black catalyst (Engelhard's fuel cell grade) with a 5 % solution of 950 e. w. Nafion[®] (Solution Technology, Inc.). Sufficient water was added to produce a viscous, but fluid mixture. The mixture was then sonicated to disperse the Pt and give a uniform

ink. This ink was brushed directly onto a Nafion* membrane, and onto a gas diffusion layer. The two electrocatalyst layers were allowed to dry separately under ambient conditions

5 Example 11

This example demonstrates the assembly of a membrane and electrode assembly (M&E) from the parts described above.

Final assembly of the M&E was accomplished by hot pressing the components to bond them together. Two gas diffusion electrodes together with a membrane were placed in a press which had been preheated to 165 °C and compressed at a pressure of 380 lbs/cm². Where Nafion® bonded electrodes had been used, the M&E was held at that temperature and pressure for approximately five minutes to "cure" the Nafion® solution to insolubility.

15 Example 12

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This example demonstrates the operation of a single cell fuel cell produced as described above.

The techniques described in examples 2, 10, and 11 were used to produce M&E's for a series of single cells. These cells were operated and the results obtained are shown by the polarization curves in Figure 15. All of these M&E's were fabricated using Nafion® 105 membrane, with 2 mg Pt/cm² thin film electrodes, and three rather disparate metallic conductors (56.5% open area expanded nickel, 31% open area expanded copper, and foamed nickel). All three polarization curves were measured with the cell operating under the same conditions, 20 to 25 °C ambient temperature, with some self-heating of the cell, air supplied by free convection, and essentially no pressure is applied to the hydrogen. As is readily apparent, all three gave essentially the same performance at current densities below 300 mA/cm², with all potential differences less than 20 mV. (The region of interest for fuel cells of this type covers the current densities from 0 to 300 mA/cm². Even at the upper end of this region, the stack will generate heat faster than it can be removed by passive means. The ensuing temperature rise will dry out the membrane, reducing performance.)

Electrodes were also produced using perforated stainless steel as the metallic conductor and yielded results significantly inferior to those shown here

Example 13

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This example demonstrates the effect of membrane selection on single cell performance.

The properties of the PEM membrane used to fabricate the M&E's is significant to their performance. Figure 16 shows the polarization curves for two M&E's operated under the same conditions, (20 to 25 °C ambient temperature, with some self-heating of the cell, air supplied by free convection, and essentially no pressure on the hydrogen), with the only difference being the membrane material used to fabricate the M&E. The cell utilizing the thinner membrane, Nafion® 112, (0.002", 0.051 mm), clearly out performs the one utilizing the thicker membrane, Nafion® 105, (0.005", 0.127 mm). From the difference in the slopes in the linear region of the polarization curves it appears that the two have different internal resistances, but the two membranes, measured under the same conditions, have very similar resistances, with the higher specific conductivity of the Nafion ® 105 offsetting its increased thickness.

The key to the difference performance is the difference in the thickness of the two membranes and how it affects water diffusion. In a typical bipolar fuel cell, both fuel and oxidizer are preferably humidified to a dew point near the operating temperature of the stack. This is done to insure that the membranes are kept fully saturated with water to maximize their conductivity. In the monopolar fuel cell, little, if any, water is supplied by the fuel, and electroosmotic drag is constantly moving the water within the membrane from the anode to the cathode, as it travels with the protons. The dew point of the air is typically 10 °C or more below the ambient temperature, which promotes evaporation from a saturated surface. If the cell has heated itself even 5 °C above ambient, it adds to the difference. Under these conditions, water is readily lost by the cathode of the fuel cell, and the membrane can dry out.

One source of water available to the membrane is the water formed at the cathode. This water can either evaporate away, or be absorbed into the membrane. It is clear that the surface of the membrane in contact with the cathode will be saturated with water, but the rest of the membrane will only have access to this water if it diffuses away from the cathode, toward the anode. The rate of diffusion increases with a reduced length of the diffusion path (i.e. thickness) and a steep concentration gradient. As a result, the thinner membrane is better humidified, and has a lower resistance during operation, for improved performance.

10 Example 14

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This example demonstrates the assembly of multi-cell modules. Assembling the multi-celled modules requires multiple press cycles, and also requires a modification of the press geometry, with the addition of a pair of blocks to act as a smaller platen than is normally available in a heated press. This permits the cell being bonded to be placed near the center of the press for maximum uniformity of both load and heat, while insuring that only one cell is pressed at a time. This arrangement is illustrated in Figure 15. Although the pins cannot be seen in this cross-section, which is intended to highlight the M&E, the two small blocks are kept in alignment by a set of four pins. These pins are firmly set in the lower block, with the upper block sliding on the pins. The use of these pins keeps the two surfaces precisely aligned through the pressing process.

The multi-cell modules produced are like the one illustrated in the crosssection shown in Figure 18. The key feature of these modules is the use of a common metal grid for the anode of one cell and the cathode of the next cell in the series. This eliminates the need for any additional current collectors or frames. This effectively reduces the number of parts needed to assemble a monopolar fuel cell module, as well as the number of electrical junctions. Each electrical junction eliminated reduces the number of potential failure points and the potential for contact resistances.

In Figure 18 the gas barrier is shown as being substantially thicker than the metal grid. While some assemblies have been produced in this manner, it is by no means a requirement of this invention. A barrier that is the same thickness as the grid is equally effective.

Example 15

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This example demonstrates the performance of a two-cell module.

Figure 19 shows a polarization curve for a two-cell module produced using M&E's produced as described in example 14, above, using Nafion® 112 membranes. The cells in this module had an active area of about 16 cm². This module produced a power output of 85 mW/cm² (for each cell) at a current density of 125 mA/cm² (0.678 Volts), the intended operating point. Although the polarization curve extends to a current density of nearly 400 mA/cm², a current density of approximately 200 mA/cm² represents the highest power that could be sustained. At higher current densities the cells gradually heated up until the membranes began to dry-out, and the performance declined.

Example 16

This example demonstrates the performance of a three-cell module.

A three-cell module was assembled as taught in example 14 and operated to determine the effect of fuel humidification on performance. Figure 20 shows the effect of fuel humidification on the performance of this three cell module. There is no significant difference at current densities below 150 mA/cm². Above 150 mA/cm² the use of humidified hydrogen produced an improvement in the module's performance. This improvement can be attributed to an improvement in the modules ability to offset the greater heating rates produced at the higher current densities, with the additional water entering with the fuel compensating for increased evaporative losses in two ways: supplying water to keep the membrane humidified, and adding extra water to remove heat by evaporation. The output of the module operating without humidification is shown in Figure 21, with the electrical and thermal contributions separated. It is clear from this figure that there is substantial heat production at higher current densities, more than can be removed by simple evaporative cooling. This is

one of the reasons for designing the system to normally operate at current densities of 125 mA/cm², or less.

Example 17

This example demonstrates the effect of fuel pressurization on the performance of a three-cell module.

All of the data in examples 15 and 16 were measured with the fuel at ambient pressure. Figure 22 shows the effect of pressurizing the fuel in the three cell system. The 0.9 psi back pressure was produced through the use of a 24 inch water column, with a slow purge maintained while operating this module and the gas exiting against the water pressure. The only region of the polarization curve where pressurization made any difference was at current densities of 200 mA/cm² and above, where a small improvement was observed. (At lower current densities, the voltages matched within ±2 mV.)

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Example 18

This example demonstrates the effect of air motion on the performance of a three-cell module.

All of the data shown in Figure 20 and in Figure 22 were obtained with air supplied solely by free convection. In order to evaluate the effect of adding forced motion and turbulence, a small fan was mounted several feet from the side of the module, and this fan used to maintain a stream of air flowing across the face of the module. Since the module is actually recessed into the face of the test fixture the stream did not strike the cathodes directly, but instead, created a large amount of turbulence over the cathode. Figure 23 shows the effect that this turbulence has on the module's performance.

It is clear that the moving air stream is reducing the output by a significant amount. There are two possible reasons for this reduction in output. The first of these is increased cooling of the module, which results in poorer kinetics, and consequently a lower potential. The second is increased drying of the membranes in the module, with the drying leading to a rise in membrane resistance, and a decline in

output. Since the drying will also lead to cooling the module, resolving these effects is difficult, but it is at least as easy to give this type of cell design too much air as it is to give it too little air.

5 Example 19

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This example illustrates the effect of using a cooling channel inside the structure of a fin-type monopolar cell structure.

Two sets of five M&Es were produced according to the procedure described in Example 14. Two frames similar to those shown in Figure 25 were attached to sheets of 3 mil (0.003", 0.76 mm) aluminum using hot melt adhesive (Bemis 3218, Bemis & Associates, Shirley, MA). To bond the frame to a sheet of aluminum, a sheet of adhesive was first trimmed to match the shape of the polymeric frame. The sheet of adhesive was placed between the frame and the metal sheet, and the whole stack placed in a heated press and bonded by pressing for five minutes at 130 °C. The metal polymer assemblies were attached to opposite sides of a sheet of convoluted aluminum, similar to the sheet depicted in Figure 27. Thermally conductive epoxy (Omegabond® 101, Omega Engineering, Stamford, CT) was used in finishing the cooling channel. The fuel cells were the attached to the polymer frames, and connected to a hydrogen supply.

A small fan was connected to the channel, and the stack was operated first with the fan in an off position and then with the fan an on position. Figure 29 is a graph showing the polarization curves for a stack operated with and without air flow from a fan. The polarization curves were obtained while operating the stack in the same conditions apart from the air flow from the fan. The stack was held at a current density of 80 mA/cm² for at least 20 minutes prior to the start of data collection. The results shown in figure 22 clearly indicate that providing an air flow through the cooling channel keeps the stack at a lower temperature resulting in an improvement of the performance. A gain of 520 mV was obtained at a current density of 100 mA/cm². When the fan is not operated, operation of the cells resulted in a heating of the stack 15 to 20° C above the ambient temperature of the laboratory (25° C), as determined by measuring the temperature at the surface of one of the cathodes. When the fan is

operated, the temperature of the system was kept between 5 and 10° C above ambient temperature. The difference in the increase in the temperature resulted in a significantly reduced rate of water loss from the cells, and improved stack potential. It is to be noted here that the fan was connected so that the air only flowed through the cooling channel and the cathodes being exposed to the same air flow in both tests.

It would be desirable to have a low pressure fuel cell that increases the overall access to air as well as air flow without the use of moving parts, allows for increased waste heat removal from the system, provides circulation of fuel through a liquid fueled system without the use of moving parts, and provides control of water at the anode and the cathode.

The invention relates to the design and fabrication of low pressure fuel cells. More particularly, the invention describes a cylindrical shape for a cell stack, a treatment for the surface of the cathodes in the stack to improve water rejection, a passive water control element to the anode compartment of the fuel cell, a sub-divided stack that produces an increased potential, and a more effective heat transfer system to improve heat rejection.

The fuel cell stack illustrated in Figures 31 (a-c) have a flat, fin-type configuration. This configuration works quite well in most applications, especially when used in conjunction with a cooling element, as illustrated in Figures 26 (a-b). However, this design may not be the most effective configuration when the air source is ambient air delivery solely by convection.

Figure 32 shows an alternate arrangement for a monopolar fuel cell 400. In this arrangement the monopolar stack 400 comprises a plurality of frusto-conical shaped stacks or cone grids. As shown a first stack 402 is positioned inside a second stack 404. The anode portion of the grid faces inward and the cathode faces outward. The first and last cone grids are essentially one-half of the other full cones to provide the connection between the anode of the top cone and the cathode of the adjacent cone and the anode of the bottom cone with a cathode. In this figure the taper of the stacks has been exaggerated for illustration purposes. Each cone has an anode 408 facing the interior of the cone, a cathode 410 facing the exterior of the cone and a proton exchange membrane 406 between the anode and the cathode. A frusto-conical

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conductive element 412 connects the anode of one stack to the cathode of the adjacent stack. This geometry allows free access to air around the full circumference of the fuel cell stack. This geometry also allows the stack to be mounted vertically on the top of the device it powers, where the stack's waste heat produces a convective air flow supplying oxygen to the cell, without any moving parts.

Figure 33 is a cross-sectional view of the cylindrical cell 400 shown in Figure 32. Membranes 406 are situated between anodes 408 and cathodes 410. Conductive elements 412 are included as part of the electrode and extend from the anode of one cell to the cathode of an adjacent cell, furnishing an electrical connection between the cells.

As described herein above and in pending US Patent application serial number 08/926,547, an internal channel may be used to remove waste heat from the interior of the fuel cell stack. While effective, a heat pipe provides greater efficiency.

Heat pipes are well known in the art and can have a cylindrical, flat, or other geometric shapes. The heat pipe is mounted in a configuration like that shown for the internal heat transfer structure, illustrated in Figure 26, and extends substantially outside the stack to furnish a greater surface area for heat dissipation.

Another aspect of the present invention relates to controlling the water at the anode. All polymers currently used for proton exchange membranes (PEM) require water for proton conduction. Water is the product of the hydrogen oxidation process, therefore it is always available, but not always in the quantities or location that are required for the efficient operation of the cell.

In an electrochemical cell, water is generated at the cathode. In a monopolar fuel cell stack the air passing over the cathode is at ambient conditions. The moisture carrying capacity of the air increases as the air is heated to the cell's operating temperature and the water eventually evaporates. Water is also removed from the cathode by back diffusion. The water concentration is highest right at the surface of the membrane, in the thin film cathode. The high concentration gradient typically causes water to diffuse down the concentration gradient, through the membrane to the anode. Back diffusion is enhanced through the use of a thin membrane, such as a 2 mil (.002", 0.054mm) thick NAFION membrane, which has a short diffusion path. A short

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diffusion path ensures that the gradient (dC/dL) is as steep as possible. If water is not removed from the anode, back diffusion will quickly act to saturate the anode and eliminate the gradient. In a typical PEM fuel cell water is continuously removed from the anode by electroosmotic drag. If the electroosmotic drag and the back diffusion are in balance, the water content in the membrane is stable.

In a hydrogen fuel cell, when a bipolar fuel cell stack is used, the water content of the membrane is stabilized through active control. This is achieved by humidification of the hydrogen fuel gas stream and the air stream before they enter the stack. Since a monopolar fuel cell stack has no means for the addition of water to either the air or the hydrogen, water control must be achieved by other means.

Figure 34 is a schematic diagram of a moisture control element 414 that can be added to the anode chamber in a hydrogen fuel cell or other fuel cell where the reactant is a gas and lack of moisture is an issue. The moisture control element 414, can be a hydrophilic foam sheet such as polyvinyl alcohol available from Rippey Corp. of Eldorado Hills, California. Other types of hydrophilic foams include but are not limited to sulfonated polystyrenes, absorbent polyacrylates, or similar hydrophilic polymers. Alternatively, a non-hydrophilic foam that has been surface treated with a surfactant to make it hydrophilic may be used, an example of this type of foam is treated polyethylene foam available from Porex Technologies of Fairburn Georgia. The moisture control element 414 serves to absorb excess water when liquid water is present at the anode and releases liquid water when the gas phase water concentration drops below saturation. The moisture control element 414 also acts as a wick to evenly distribute water within each anode compartment.

Figure 34 shows a typical membrane and electrode assembly (M&E) 416 with a polymer flow grid 418 between the M&E 416 and the moisture control element. The polymer flow grid 418 is provided to promote fuel gas flow to the anode surface and provide some stability to the structure. An aluminum or other type of gas barrier 420 may be provided adjacent to the moisture control element. The gas barrier in addition to inhibiting gas flow, it can provide some heat removal. Alternatively, a heat pipe (discussed above) can be positioned adjacent to the moisture control element 414

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The moisture control element 414 may also be used to improve the distribution area of methanol in a direct methanol fuel cell (DMFC) fabricated using the low pressure stacks described herein.

Another aspect of the present invention provides an improved stack design resulting in greater power output and efficiency for a DMFC. A stack assembly for a 150 Watt battery charger based on monopolar direct methanol fuel cells is illustrated in Figure 35. As shown, there are four fin-type stacks 422, 424, 426, 428, with fourteen cells on each side of each fin, for a total of 112 cells. The four fins are connected in a combination series-parallel arrangement, with each fin paired in parallel with an adjacent fin and the pair of parallel fins are connected in series with the other pair of fins. CO₂ vents 430 are provided at the top of the stack to collect CO₂ generated by the electrodes. A fuel feed manifold 434 is in flow communication with bottom of the stacks for delivering liquid fuel such as methanol.

All four fins are oriented with space between each fin so that air is easily lifted through the structure by convection. At the same time as the air moves up along the stacks, the water, directed by the leaders 429, runs down the edges of the fins, to be collected at the bottom of the stack.

Air flow is adjusted for climate conditions with three manually adjusted dampers, illustrated in Figure 36. The damper or exhaust vent 436 at the top acts to restrict air flow under any conditions to prevent excessive air flow from over cooling the stack. The other two dampers are used to adjust for extremes of temperature. When the bottom damper 438 is open, air is admitted directly to the fins for operating under mild to warm conditions. Under cool to cold conditions, the lower damper is at least partially closed, and the upper damper 440 opened. This causes the air to enter at the top of the structure. The air is then prewarmed, and the stack is slightly cooled, as the air is siphoned down the outside of the stack before it is pulled in, and up the inside of the stack by convective heating.

This stack design can be used with methanol fuel. Initially, the methanol is added as a concentrated solution, where the stack itself operates with a more dilute solution. The exact composition of the solution in the fuel loop is a function of the membrane used to fabricate the stack. Although stacks equipped with NAFION

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membranes usually operate at less than one molar concentration, improved membranes with better resistance to methanol cross over may permit the use of higher concentrations. Variations that may result from the substitution of a superior membrane material are considered to be within the scope of this application.

Like the air, the solution within the stack changes temperature on passing through the stack, and is lifted by convection. In addition, the generation of carbon dioxide at the anode produces buoyant bubbles which act as a pump to move the fuel solution through the loop. Each of the four fins in this power supply generate 306 mL of $\rm CO_2$ per minute based on the molar volume of the gas as measured under standard conditions.

Figure 37 is a schematic view of the fuel loop 442, including a recycle loop 443. The CO₂ bubbles 444 generated at the anode direct the flow of the methanol through the loop. The CO₂ is collected at the CO₂ vent 430. An electrochemical methanol sensor 446 is positioned near the end of the loop above the point of methanol delivery 448. As the main fuel loop circulates, its composition is constantly monitored by the electrochemical methanol sensor 446. When the methanol content drops below a predetermined level, additional methanol solution is injected into the loop. When the methanol concentration rises above a predetermined level, recovered water is injected into the loop at injection point 450 to dilute the fuel stream.

In a completely integrated monopolar DMFC battery charger, as described here, the number of moving parts is minimized, and thus the amount of power consumed by the charger as a whole is also minimized.

Another embodiment of the present invention includes providing a thin layer of water repellent compound such as polytetraflouroethylene (PTFE) on the cathode surface. This water repellent layer can be applied as a coating through the use of a suspension or solution of the water repellent compound, or applied as a thin film. The presence of the water repellent layer aids in the removal of water produced at the cathode. As water vapor condenses on the cooler outer surface of the monopolar structure, the droplets quickly run off without blocking any of the active surface area of the cathode. As shown in Figure 38, the cathode 452 has a PTFE layer 454 on its

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surface. The membrane 456 and anode 458 have conductive elements 460 embedded therein as described previously.

Also disclosed here is a geometry intended to increase the voltage of a monopolar fuel cell stack. In all of the designs shown previously, each side of each fin had one series of cells. In the design shown in Figure 39 three rows of cells are all arranged on the same surface. Each set of five cells is individually fabricated, and all three are attached to a common support frame, with the individual groups of cells kept electrically isolated from each other excSept at the connection points at the two ends. A cathode 462 is at one end of the top row. At the end of the top row, the anode 464 is connected to the cathode 466 at the end of the second row. The opposite end of the second row, an anode 468 is connected to a cathode 470 on the third row. The path followed by the current flow through this portion of the stack is illustrated in Figure -40. While the use of this arrangement leads to smaller cells, the increased number of cells in series produces a substantial increase in voltage.

Example 20

The following example illustrates the performance of a stack without hydrophilic foam to the anode compartment.

A fuel cell stack was fabricated consisting of four fins like the one shown in Figure 31. Each of these fins had five cells on each of 2 opposing faces, with a cooling channel located between the adjacent fins. Each individual cell had an active area of 20 cm². All of the cells on each fin were electrically connected in series to produce a potential in excess of 6 Volts. The fins were interconnected with two fins in series and the two series of fins connected in parallel to produce a stack capable of delivering 4 Amps of power at a potential in excess of 12 Volts.

The performance of this nominally 50 Watt power supply is illustrated by a standard i-V curve in Figure 41. This same data is replotted as power vs. current in Figure 42. This stack could only achieve full power operation for short periods of time and exhibited evidence of the membranes drying when operated for extended periods at any current in excess of 2 Amps. This data represents the performance of a 50W stack design, without the inclusion of the hydrophilic water control sponges in the anode gas chambers

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Example 21.

This example illustrates the benefits of adding a hydrophilic foam to the anode compartment.

A second fuel cell stack was fabricated like the one described in Example 1. This stack included a hydrophilic foam element for water control like that shown in Figure 34.

Figure 43 and Figure 44 show the operating data for the 50 Watt fuel cell stack including the hydrophilic foam elements. This stack had superior performance at higher currents and could maintain full power for extended periods.

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While the foregoing is directed to the preferred embodiment of the present invention, other and further embodiments of the invention may be devices without departing from the basic scope thereof, and the scope thereof is determined by the claims which follow.

What is claimed is:

- An apparatus comprising an array of cells; each cell comprising:
- 2 (a) a membrane:
- 3 (b) an anode comprising a conductive element; and
- 4 (c) a cathode comprising a conductive element:
- 5 wherein the cathode of one cell shares its conductive element with the anode of
- 6 an adjacent cell; the sharing of conductive elements establishing an electrical
- 7 connection between the cells in the array.
- 1 2. The apparatus of claim 1 wherein the cells are electrochemical cells suitable for
- 2 the direct production of oxidized and/or reduced products through simultaneous
- 3 electrochemical oxidation and reduction of a compound or a mixture of compounds by
- 4 application of an electrical current.
- The apparatus of claim 2 wherein the electrochemical cells are suitable for the
- 2 production of hydrogen and oxygen by hydrolysis of water.
- 1 4. A method for producing hydrogen and oxygen by electrolysis of water in the
- 2 apparatus of claim 3.
- The apparatus of claim 1 wherein the cells are fuel cells suitable for the direct
- 2 generation of electricity by oxidation of a fuel and reduction of an oxidizer
- The apparatus of claim 5 wherein the fuel is a gas.
- The apparatus of claim 6 wherein the fuel is hydrogen gas.
- 1 8. The apparatus of claim 6 wherein the fuel is a gas mixture comprising
- 2 hydrogen.

- The apparatus of claim 5 wherein the fuel is a liquid.
- 1 10. The apparatus of claim 9 wherein the fuel is an alcohol, an ether or a mixture
- 2 thereof; said mixture being either neat, or in aqueous solution.
- 1 11. The apparatus of claim 10 wherein the fuel is selected from methanol.
- 2 dimethoxymethane, trimethoxymethane, and mixtures thereof.
- 1 12. The apparatus of claim 5 wherein the oxidizer is air.
- 1 13. The apparatus of claim 12 wherein the air is provided by unassisted
- 2 convection.
- 1 14. The apparatus of claim 12 further comprising an air mover for assisted
- 2 convention.
- 1 15. The apparatus of claim 14 wherein the air is provided at a pressure less than
- 2 0.5 psi above ambient pressure.
- 1 16. The apparatus of claim 14 wherein the air mover is a fan.
- A low pressure gas electrode comprising
- 2 (a) an electrically conducting supporting substrate; wherein said substrate
 - comprises at least two opposite faces; a plurality of openings allowing
- 4 passage from one face to an opposite face; said openings encompassing
- 5 between 10% and 90 % of the total area of the substrate; and
- 6 (b) an electrically conductive porous material bound to the substrate comprised
- 7 of a conductive component and a binding component; and
- 8 (c) an electrocatalyst layer bound to the substrate.
- 1 18. The electrode of claim 17 wherein the binding component is electrically

- conductive.
- 1 19. The electrode of claim 17 wherein the binding component is electrically non
- conductive.
- 1 20. The electrode of claim 17 wherein the passage openings represent between 20
- 2 % and 80 % of the total area of the substrate.
- 1 21. The electrode of claim 17 wherein the passage openings represent between
- 2 30% and 75 % of the total area of the substrate.
- 1 22. The electrode of claim 17 wherein the passage openings represent between 24
- 2 % and 70 % of the total area of the substrate.
- 1 23. The electrode of claim 17 wherein the electrically conductive porous material is
- 2 a mixture comprising a powdered component and a fibrous component.
- 1 24. The electrode of claim 23 wherein the powdered component comprises carbon.
- 1 25. The electrode of claim 24 wherein the fibrous component comprises carbon.
- The electrode of claim 23 wherein the fibrous component comprises carbon.
- 1 27. The electrode of claim 25 wherein the ratio of powdered carbon to fibrous
- 2 carbon is between 1:1 and 5:1.
- The electrode of claim 27 wherein the ratio is between 2:1 and 3:1.
- 1 29. The electrode of claim 23 wherein the powdered component is a metal.
- 1 30. The electrode of claim 19 wherein the binding component is a polymer.

- 1 31. The electrode of claim 30 wherein the polymer is at least partially fluorinated.
- 1 32. The electrode of claim 31 wherein the polymer is a fully fluorinated
- 2 thermoplastic.
- 1 33. The electrode of claim 32 wherein the polymer is polytetrafluoroethylene.
- 1 34. The electrode of claim 17 wherein the ratio of the binding component to the
- 2 conductive component is between 2:1 and 1:20; preferably between 1:1 and 1:8; and
- 3 more preferably between 1:1 and 1:3.
- 1 35. The electrode of claim 17 wherein the electrically conductive supporting
- 2 substrate comprises a metal.
- 1 36. The electrode of claim 35 wherein the substrate is a sheet of expanded metal.
- 1 37. The electrode of claim 35 wherein the substrate is a piece of woven wire fabric.
- 1 38. The electrode of claim 37 wherein the substrate is a screen.
- 1 39. The electrode of claim 35 wherein the electrically conductive supporting
- 2 substrate is a perforated metal sheet.
- 1 40. The electrode of claim17 wherein an electrocatalyst layer is deposited on one
- 2 face of said substrate.
- 3 41. An electrochemical device having an electrochemical cell comprising the
- 4 electrode of claim 17.
- The electrochemical device of claim 41 wherein the electrochemical cell is an
- 2 electrolytic cell.

- 1 43. The electrochemical device of claim 42 wherein the electrolytic cell is an
- electrolyzer for the generation of hydrogen and oxygen from water.
- 44. The electrochemical device of claim 41 wherein the electrochemical cell is a
- 2 fuel cell for the generation of electricity.
- 1 45. The electrochemical device of claim 44 wherein the cells are connected in a
- 2 series to form an assembly with increased electrical potential.
- 1 46. The electrochemical device of claim 45 wherein the electrochemical cells are
- 2 arranged in a bipolar configuration.
- 1 47. The electrochemical device of claim 45 wherein the electrochemical cells are
- 2 arranged in a monopolar configuration.
- 1 48. A dual electrode structure comprising a first electrode and a second electrode,
- 2 wherein the electrodes comprise:
- 3 (a) an electrically conducting supporting substrate; wherein said substrate
- 4 comprises at least two opposite faces; a plurality of openings allowing
- 5 passage from one face to an opposite face; said openings encompassing
 - between 10% and 90 % of the total area of the substrate:
- 7 (b) an electrically conductive porous material bound to the substrate comprised
- 8 of a conductive component and a binding component;
- (c) an electrocatalyst layer bound to the substrate;
- and wherein the two electrodes share the same electrically conducting
- 11 substrate.
 - 1 49. The dual electrode structure of claim 48 further comprising a barrier to gas
- 2 passage; wherein the barrier is formed between the two electrodes by sealing all of the
- 3 open spaces of the conductive substrate; and wherein the electrodes having their
- 4 electrocatalyst layers bound to opposite sides of the electrically conducting substrate.

- 50. A fuel cell stack comprising a plurality of dual electrode structures; each dual
 electrode structure comprising a first electrode and a second electrode, wherein the
 electrodes comprise:
- (a) an electrically conducting supporting substrate; wherein said substrate
 comprises at least two opposite faces; a plurality of openings allowing
 passage from one face to an opposite face; said openings encompassing
 between 10% and 90 % of the total area of the substrate:
- (b) an electrically conductive porous material bound to the substrate comprised
 of a conductive component and a binding component;
- (c) an electrocatalyst layer bound to the substrate;
 - and wherein the two electrodes share the same electrically conducting substrate; and wherein the first electrode forms the anode of a first fuel cell and the second electrode forms the cathode of an adjacent fuel cell; the fuel cells being connected in series to produce a potential larger than the potential of each individual fuel cell.
- 1 51. An apparatus comprising
- 2 (a) a frame having two opposing faces, and
 - (b) a set of first and second fuel cell stacks of claim 50, wherein the first fuel cell stack is attached to one face of the frame and the second fuel cell stack is attached to the opposite face of the frame; and wherein the set of stacks and the frame define a volume for internal storage of a reactant to be consumed by operation of the fuel cells;
- 8 (c) means to supply said internally stored reactant; and
- (d) means to remove undesirable waste accumulated by operation of the fuel
 cell stacks.
- 1 52. An apparatus comprising:
- (a) a frame having two opposing faces;
- 3 (b) a barrier, and
- 4 (c) a fuel cell stacks of claim 50; wherein the fuel cell stack is attached to one

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- face of the frame and the barrier is attached to the opposite face of the frame; and wherein the stack, the barrier and the frame define a volume for internal storage of a reactant to be consumed by operation of the fuel cells;
- 8 (d) means to supply said internally stored reactant; and
- (e) means to remove undesirable waste accumulated by operation of the fuel
 cell stacks.
- 1 53. The apparatus of claim 52 manufactured from flexible materials, and wherein
- 2 the barrier has a surface with a curved shape; the apparatus being mounted on the
- 3 surface of the barrier and adopting the shape of the surface.
- 1 54. The apparatus of claim 52 further comprising an enclosed channel for the flow
- 2 of a coolant to dissipate heat generated by operating the apparatus; and wherein the
- 3 barrier provides one surface of the channel.
- The apparatus of claim 54 wherein the channel is surrounded by at least one set
- of electrochemical cells.
- 1 56. The apparatus of claim 54 further comprising a heat transfer element
- 2 positioned within the channel.
- 1 57. The apparatus of claim 54 further comprising means for improving the flow of
- 2 the coolant through the channel.
- 1 58. The apparatus of claim 52 wherein the barrier has a portion that extends
- 2 beyond the cells; said portion allowing the dissipation of the heat generated while
- 3 operating the apparatus.
- 1 59. The apparatus of claim 58 wherein the barrier's portion extending beyond the
- 2 fuel cells is bent to extend perpendicularly to the plane of the cells and providing a
- 3 protection to portions of the electrodes that are exposed to the atmosphere.

- 1 60. The apparatus of claim 59 wherein the barrier is perforated and further bent to
- 2 form a cover shield to protect portions of the electrodes that are exposed to the
- 3 atmosphere.
- A method of making a porous electrode comprised of:
- 2 (a) mixing high surface area carbon powder with a polymer electrolyte
- 3 suspension to form a mixture;
- 4 (b) sonicating said mixture to reach complete dispersion and form a paste; and
- 5 spreading said paste on a foil grid to form a gas diffusion electrode.
- An apparatus comprising at least two arrays of cells; each cell comprising:
- (a) a membrane;
- 3 (b) an anode comprising a conductive element; and
- 4 (c) a cathode comprising a conductive element;
- 5 wherein the cathode of one cell shares its conductive element with the anode of
- an adjacent cell; the sharing of conductive elements establishing an electrical
- 7 connection between the cells in the array.
- 8 wherein arrays have a generally conical shape.
- 1 63. The apparatus of claim 62 wherein one of the arrays is adapted to surround a
- 2 portion of the other array in a substantially circular pattern.
- 1 64. The apparatus of claim 62, wherein the cells are electrochemical cells suitable
- 2 for the direct production of oxidized and/or reduced products through simultaneous
- 3 electrochemical oxidation and reduction of a compound or a mixture of compounds by
- 4 application of an electrical current.
- 1 65. The apparatus of claim 62 further comprising a reactant chamber formed
- 2 between the anode surfaces of adjacent arrays.
- 1 66. The apparatus of claim 62 wherein the conductive elements form a generally

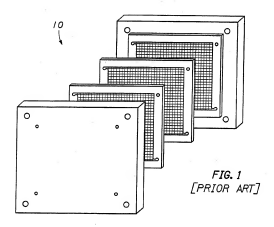
- 2 conical shape.
- 1 67. An array of electronically conducting cells, wherein each cell comprises:
- 2 a membrane electrode assembly having an anode comprising a conductive
- 3 element and a cathode comprising a conductive element;
- 4 wherein the cathode of one cell shares its conductive element with the anode of
- 5 an adjacent cell so that an electrical connection is established between the cells in the
- 6 array; and
- 7 a moisture control element adjacent to the anode.
- 1 68. The array of claim 67 wherein the moisture control element is a hydrophilic
- 2 foam.
- 1 69. The array of claim 67 further comprising a support element between the
- 2 moisture control element and the anode.
- 1 70. The array of claim 67 wherein the moisture control element is adapted to
- 2 absorb liquid water when a water concentration exceeds saturation.
- 1 71. The array of claim 67 wherein moisture control element is adapted to release
- 2 liquid water when the gaseous water concentration drops below saturation.
- 1 72. The array of claim 67 wherein the moisture control element is selected from
- 2 polyvinyl alcohols, polyacrylates, sulfonated polystyrenes and mixtures thereof.
- 1 73. The array of claim 68 wherein the support element is a polymer flow grid.
- 1 74. An array of electronically conducting cells, wherein each cell comprises
- a membrane electrode assembly having an anode comprising a conductive
- 3 element and a cathode comprising a conductive element;
- 4 wherein the cathode of one cell shares its conductive element with the anode of

- an adjacent cell so that an electrical connection is established between the cells in the 5 6 array: and
- 7 a moisture control element adjacent to the anode for absorbing liquid water and releasing liquid water when gaseous water concentration is below saturation. 8
- The array of claim 74 wherein the moisture control element is selected from polyvinyl alcohols, polyacrylates, sulfonated polystyrenes and mixtures thereof. 2
 - 76. An apparatus comprising:

- an array of monopolar cells, each cell comprising a membrane and electrode 2 assembly and first and second electrically conducting frames contacting opposing faces 3 of the membrane and electrode assembly and electrically connecting the first electrically conducting frame of a first cell to the second electrically conducting frame 5 of an adjacent cell-
- 7 a fuel gas chamber defined by the array and an electrically insulating member sealed to the edge of the array; 8
- 9 a fuel gas inlet port into the chamber, and
- a waste gas outlet out of the chamber. 10
- 1 The apparatus of claim 76 further comprising a housing surrounding the array of monopolar cells, wherein the housing has a plurality of vents for controlling the air 2 flow through the array by convection.
- 3
- 1 78 The apparatus of claim76 wherein the array comprises a plurality of fins
- supporting the cells, wherein pairs of adjacent fins are electrically connected in parallel 2
- 3 with one another and those pairs that are connected in parallel, are connected in series
- with an adjacent pairs of fins.
- 1 79 The apparatus of claim 78, wherein each fin is positioned parallel to an adjacent
- fin. 2
- 3

- 80. An array of electronically conducting cells, wherein each cell comprises:
- a membrane electrode assembly having an anode comprising a conductive

 element and a cathode comprising a conductive element;
- 4 wherein the cathode of one cell shares its conductive element with the anode of
- 5 an adjacent cell so that an electrical connection is established between the cells in the
- 6 array; and
- 7 a moisture control element adjacent to the cathode.
- 1 81. The array of claim 80, wherein the moisture control element comprises a layer
- 2 of water repellent material.



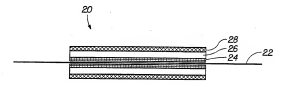


FIG. 2 [PRIOR ART]

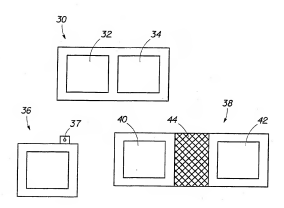


FIG. 3

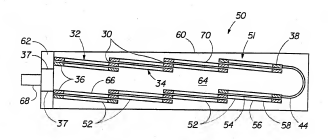


FIG. 4

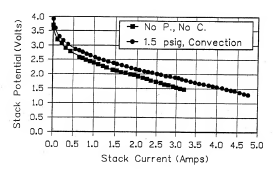


FIG. 5

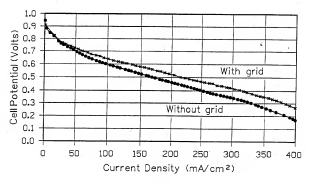


FIG. 7

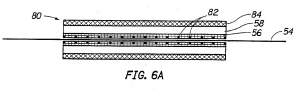
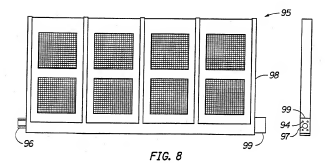




FIG. 6B



FIG. 6C



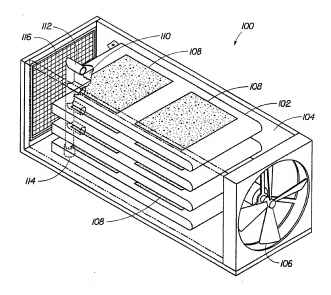
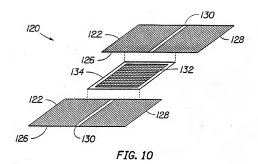
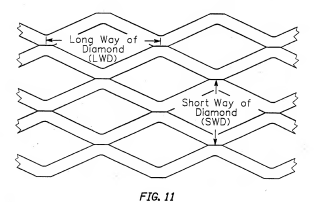
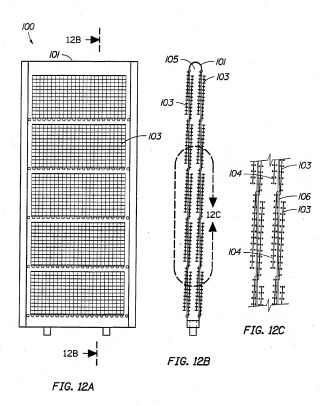


FIG. 9







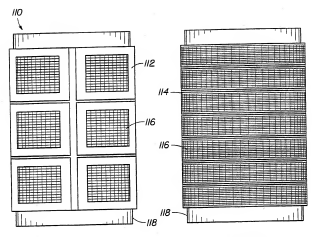


FIG. 13A

FIG. 13B

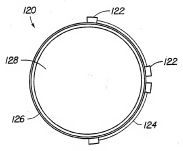


FIG. 14

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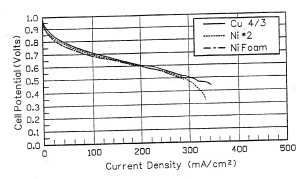


FIG. 15

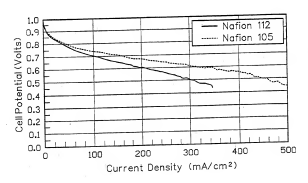
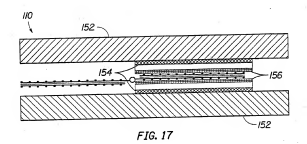


FIG. 16



164 162 164 168 168 FIG. 18

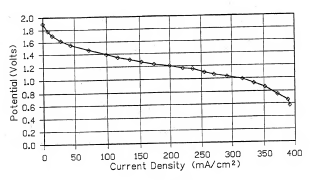


FIG. 19

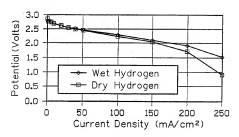


FIG. 20

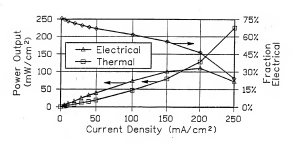


FIG. 21

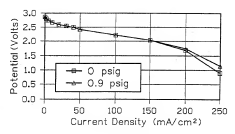


FIG. 22

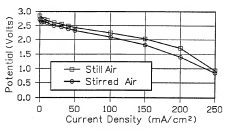


FIG. 23

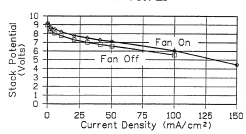


FIG. 24
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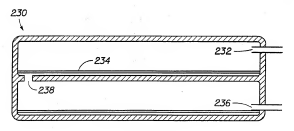
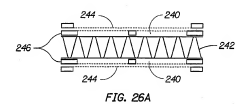
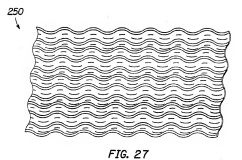


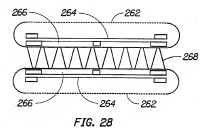
FIG. 25

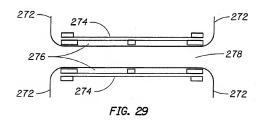


244 244

FIG. 26B







80 58 56 5

FIG. 30A

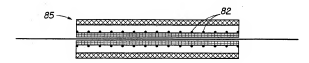


FIG. 30B

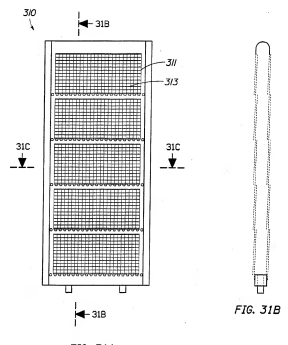


FIG. 31A



FIG. 31C

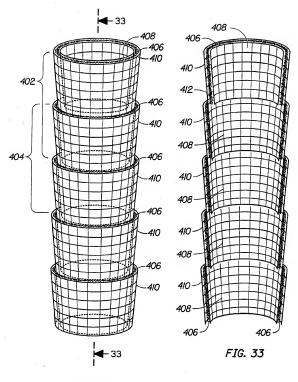
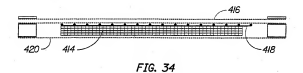


FIG. 32



430 432 430 432 429 428 428

FIG. 35



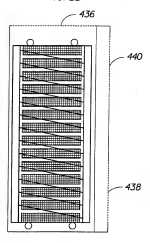


FIG. 36

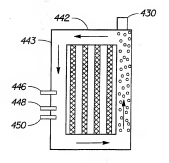
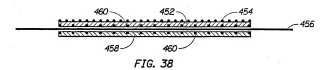


FIG. 37



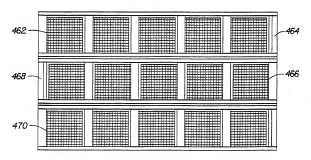


FIG. 39

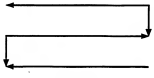


FIG. 40

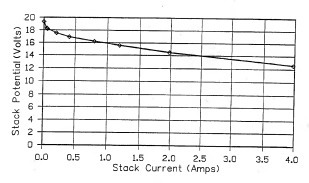


FIG. 41

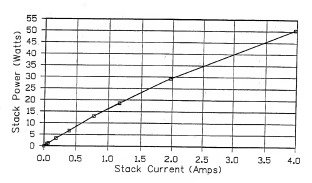


FIG. 42

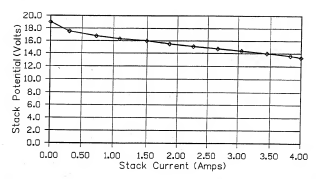


FIG. 43

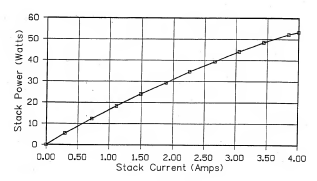
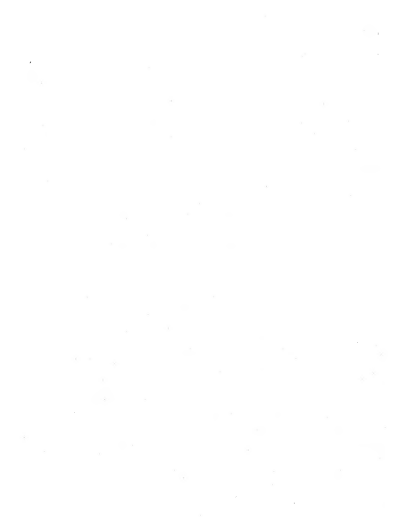


FIG. 44



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(51) International Patent Classification ⁶ :		(11) International Publication Number: WO 99/3446
H01M 8/24, 8/04, 4/86, 4/96, 4/88, C25B 9/00, 11/03	A3	(43) International Publication Date: 8 July 1999 (08.07.9
(21) International Application Number: PCT/US (22) International Filing Date: 10 September 1998 (30) Priority Data: 08/926.547 10 September 1997 (10.09.5 (63) Related by Continuation (CON) or Continuation-li (CIP) to Earlier Application US Not furnis Filed on Not furnis Filed on Not furnish Filed on Priority (10.09.5) No.	(10.09.9 n-Part hed (Clifumishe NNTEClifumishe Lifumishe NNTEClifumishe NNTEClif	BY, CA, CH, CN, CU, CZ, DE, DK, EE, BS, FI, GB, Gi GH, GM, HUJ, DH, IS, JP, FE, KG, KP, KR, KZ, L LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, M NO, NC, PL, PT, RO, RU, SD, SE, SG, SI, SK, JL, TT, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARPIO pate (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian pate (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European pate (AT, BE, CH, CY, DE, DK, ES, FI, RG, GR, RI, ET, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, C M, GA, GN, GW, ML, MR, NE, SN, TD, TG). Published With international search report. [88] Date of publication of the international search report: 11 November 1999 (11.11.5
as well as a simpler design that permits the size of the the present design, an adjacent anode and cathode pair an serving to conduct the current from one cell to the adjacet fuels or alternatively directly with liquid fuels, such as me ermits the storage of more energy in less volume while at	vention system e fabric nt one. thanol, the san	has a reduced number of component parts to reduce fabrication cost to be reduced at the same time as performance is being improved. Indeed using a common conductive element, with that conductive element. This produces a small and simple system suitable for operating with grimenhoxymethance, or trimentoxymethane. The use of these liquid the time eliminating the need for handling compressed gaines which furthlessing of this invention can be further increased by adding a passage for the suitable of the suita
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INTERNATIONAL SEARCH REPORT

ir stional Application No

PCT/US 98/19221 A. CLASSIFICATION OF SUBJECT MATTER
IPC 6 H01M8/24 H01M8/04 H01M4/86 H01M4/96 H01M4/88 C25B9/00 C25B11/03 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 6 HO1M C25B Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base end, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category o Citetion of document, with indication, where appropriate, of the relevant passages Relevant to claim No US 5 364 711 A (YAMADA SHUJI ET AL) 48-60. 15 November 1994 (1994-11-15) 62-66 claims 1-20 US 4 596 648 A (SWEENEY CHARLES T) 1-4 24 June 1986 (1986-06-24) claims 1-23 X US 4 235 693 A (MCCARTNEY JOSEPH F ET AL) 1 - 8.7625 November 1980 (1980-11-25) claims 1-17 X US 5 053 375 A (RAO BHASKARA M L) 17-19. 1 October 1991 (1991-10-01) 23-27 claims 1-14 -/--X Further documents are listed in the continuation of box C. X Patent family members are listed in annex. Special categories of cited documents: "T" later document published after the international filing date or priority date and not in conflict with the epplication but "A" document defining the general state of the lart which is not considered to be of particular relevence cited to understand the principle or theory underlying the "E" earlier document but published on or after the Internetional "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone filing date "L" document which may throw doubte on priority claim(e) or which is cited to establish the publication date of enother citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled in the art. "P" document published prior to the international filling date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 2 September 1999 10/09/1999 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijewijk Tel. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3015 Battistig, M

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C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT Category . Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. US 3 854 994 A (KOHLING A ET AL) 61 χ 17 December 1974 (1974-12-17) claims 1-19 23-40 Α PATENT ABSTRACTS OF JAPAN 62 X vol. 018, no. 626 (E-1636), 29 November 1994 (1994-11-29) & JP 06 243879 A (FUJI ELECTRIC CO LTD). 2 September 1994 (1994-09-02) abstract US 4 364 805 A (ROGERS DOUGLAS K) 17-40 Α 21 December 1982 (1982-12-21) claims 1-6 US 5 599 638 A (SURAMPUDI SUBBARAO ET AL) 1-16, 4 February 1997 (1997-02-04) 41-47. 50-60, 62-80 claims 1-49 WO 94 25991 A (AER ENERGY RESOURCES INC) 77-81 10 November 1994 (1994-11-10) claims 1-32

2

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INTERNATIONAL SEARCH REPORT Information on patent family members

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	US	4596648	A	24-06-1986	NON	:			
	US	4235693	A	25-11-1980	NON	·			
	US	5053375	Α	01-10-1991	AU CA EP MX WO	7242591 A 2072088 A 0510107 A 166747 B 9111033 A	05-08-1991 09-07-1991 28-10-1992 01-02-1993 25-07-1991		
	US	3854994	A	17-12-1974	DE	2208632 A	06-09-1973		
	JP	06243879	A	02-09-1994	NON	:			
	US	4364805	Α	21-12-1982	NONE				
	US	5599638	A	04-02-1997	US	5773162 A	30-06-1998		
	WO	9425991	A	10-11-1994	AT AU CA DE DE EP JP US US	146306 T 6823294 A 2161668 A 69401130 D 69401130 T 0696384 A 8511896 T 5560999 A 5721064 A	15-12-1996 21-11-1994 10-11-1994 23-01-1997 28-05-1997 14-02-1996 10-12-1996 01-10-1996 24-02-1998		